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- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

### IMAGES ARE BEST AVAILABLE COPY.

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=> d que stat 14
           4360 SEA FILE=HCAPLUS ABB=ON (RSV? OR ?RESPIRATOR?(W)?SYNCYTIAL?(W)
                ?VIRUS?)
            187 SEA FILE=HCAPLUS ABB=ON L1 AND ?IMMUNOGEN?
L2
            138 SEA FILE=HCAPLUS ABB=ON L2 AND (F OR G OR M OR SH OR NS1? OR
L3
                NS2? OR P)
             13 SEA FILE=HCAPLUS ABB=ON L3 AND M2?
=> d ibib abs 14 1-13
    ANSWER 1 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN
                         2003:625764 HCAPLUS
ACCESSION NUMBER:
                         Evaluation of recombinant respiratory
TITLE:
                         syncytial virus gene deletion
                         mutants in African green monkeys for their potential
                         as live attenuated vaccine candidates
                         Jin, Hong; Cheng, Xing; Traina-Dorge, Vicki L.; Park,
AUTHOR(S):
                         Hyun Jung; Zhou, Helen; Soike, Ken; Kemble, George
                         MedImmune Vaccines Inc., 297 North Bernardo Avenue,
CORPORATE SOURCE:
                         Mountain View, CA, 94043, USA
                         Vaccine (2003), 21(25-26), 3647-3652
SOURCE:
                         CODEN: VACCDE; ISSN: 0264-410X
                         Elsevier Science Ltd.
PUBLISHER:
DOCUMENT TYPE:
                         Journal
                         English
LANGUAGE:
     Towards the goal of developing live attenuated respiratory
     syncytial virus (RSV) vaccines to prevent
     severe respiratory tract infections caused by respiratory
     syncytial virus, recombinant RSV contg. a
     deletion of single or multiple NS1, NS2, SH
     and M2-2 genes have been generated. In this study,
     recombinants, rA2.DELTA.M2-2, rA2.DELTA.NS2,
     rA2.DELTA.NS1NS2, rA2.DELTA.SHNS2, rA2.DELTA.M2-2NS2
    were evaluated in African green monkeys (AGMs) for their infectivity,
     immunogenicity and protection against wild type (wt) RSV
     challenge. Replication of rA2.DELTA.NS2 and rA2.DELTA.SHNS2 was
    not attenuated in either the upper or the lower respiratory tracts of
           On the other hands, rA2.DELTA.NS1NS2 was over-attenuated;
     it did not replicate in the respiratory tracts of the infected monkeys and
     did not provide sufficient protection against wild type RSV
     challenge. rA2.DELTA.M2-2NS2 was slightly more attenuated than
    rA2.DELTA.M2-2 and provided partial protection against wt
    RSV challenge. rA2.DELTA.M2-2, and possibly rA2.DELTA.
    M2-2NS2, exhibited the attenuated but protective phenotypes in the
    monkeys that could be further evaluated as potential live attenuated
    RSV vaccine candidates in the clin. studies.
    ANSWER 2 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN
                         2003:511167 HCAPLUS
ACCESSION NUMBER:
                         139:51610
DOCUMENT NUMBER:
                         Immunogenic compositions comprising an
TITLE:
                         antigen and a purified {\bf M} protein from
                         respiratory syncytial virus
                         Barber, Brian; Cates, George; Parrington, Mark;
INVENTOR(S):
                         Sambhara, Suryprakash
                         Aventis Pasteur Limited, Can.
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 27 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                        English
LANGUAGE:
```

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

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APPLICATION NO. DATE
     PATENT NO.
                     KIND
                            DATE
                                        WO 2002-CA1953 20021218
                     A1
                            20030703
     WO 2003053464
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD,
             RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
             CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,
             MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                        US 2001-341422P P 20011220
    Methods and compns. for enhancing an immune response to an antigen in a
    host are provided. Immunogenic compn. comprising an antigen and
     an amt. of purified M protein from respiratory
     syncytial virus are provided in a pre-selected amt. to
     provide an enhanced immune response to said antigen in a host having a
    pre-existing respiratory syncytial virus
    M-specific immune response. The antigen can be an antigen from
     respiratory syncytial virus.
REFERENCE COUNT:
                         4
                               THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 3 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN
L4
                         2002:869921 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         138:135888
TITLE:
                         Recombinant Respiratory syncytial
                         virus with the G and F
                         genes shifted to the promoter-proximal positions
                         Krempl, Christine; Murphy, Brian R.; Collins, Peter L.
AUTHOR(S):
                         Laboratory of Infectious Diseases, National Institute
CORPORATE SOURCE:
                         of Allergy and Infectious Diseases, Bethesda, MD,
                         20892-8007, USA
                         Journal of Virology (2002), 76(23), 11931-11942
SOURCE:
                         CODEN: JOVIAM; ISSN: 0022-538X
PUBLISHER:
                         American Society for Microbiology
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
     The genome of human respiratory syncytial
AB
     virus (RSV) encodes 10 mRNAs and 11 proteins in the
     order 3'-NS1-NS2-N-P-M-SH
     -G-F-M2-1/M2-2-L-5'. The
     G and F glycoproteins are the major RSV
     neutralization and protective antigens.
                                               It seems likely that a high level
     of expression of G and F would be desirable for a live
     RSV vaccine. For mononegaviruses, the gene order is a major
     factor controlling the level of mRNA and protein expression due to the
     polar gradient of sequential transcription. In order to increase the
     expression of G and F, recombinant RSVs
     based on strain A2 were constructed in which the G or F
     gene was shifted from the sixth or seventh position (in a genome lacking
     the SH gene), resp., to the first position (rRSV-G1/.DELTA.
     SH and rRSV-F1/.DELTA.SH, resp.). Another virus was
     made in which G and F were shifted together to the
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first and second positions, resp. (rRSV-G1F2/.DELTA.SH). Shifting one or two genes to the promoter-proximal position resulted in increased mRNA and protein expression of the shifted genes, with G and F expression increased up to 2.4-and 7.8-fold, resp., at the mRNA level and approx. 2.5-fold at the protein level, compared to the parental virus. Interestingly, the transcription of downstream genes was not greatly affected even though shifting G or F, or G and F together, had the consequence of moving the block of genes NS1-NS2-N-P-M-( G) one or two positions further from the promoter. The efficiency of replication of the gene shift viruses in vitro was increased up to 10-fold. However, their efficiency of replication in the lower respiratory tracts of mice was statistically indistinguishable from that of the parental virus. In the upper respiratory tract, replication was slightly reduced on some days for viruses in which G was in the first position. The magnitude of the G-specific antibody response to the gene shift viruses was similar to that to the parental virus, whereas the F-specific response was increased up to fourfold, although this was not reflected in an increase of the neutralizing activity. Thus, shifting the G and F genes to the promoter-proximal position increased virus replication in vitro, had little effect on replication in the mouse, and increased the antigen-specific immunogenicity of the virus beyond that of parental RSV.

REFERENCE COUNT: 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 4 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER:

2002:142851 HCAPLUS

DOCUMENT NUMBER:

136:215388

TITLE:

Immunogenic hepatitis B nucleocapsid protein

(HBc) chimeric particles having enhanced stability

INVENTOR(S):

Birkett, Ashley J. Apovia, Inc., USA

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 290 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

P.F	TENT	NO.		KI	ND	DATE		A.	PPLI	CATI	ои ис	٥.	DATE			
	2002							W	0 20	01-U	S417	59	2001	0816		
WC		ΑE,	AG,	AL,	ΑU,	BA,	BB,		-				CR,			
													KR, SG,			
	RW:	•	•	•	•		•				•	•	TJ, AT,		CH,	CY,
													PT, SN,			BF,
	2003	1387	69	Α	1	2003	0724	Ü	S 20	01-9	3091	5	2001	0815		
	7 2001 2 1333	857		A	2	2003	0813	E.	P 20	01-9	6461	5		0816		
	R:					DK, FI,					LI,	LU,	NL,	SE,	MC,	PT,
PRIORIT	Y APF		•		·		ŕ	US 2	000-	2258			2000			
													2001			

WO 2001-US41759 W 20010816

AB A chimeric, carboxy-terminal truncated hepatitis B virus nucleocapsid protein (core protein or HBc) is disclosed that is engineered for both enhanced stability of self-assembled particles and the display of an immunogenic epitope. The immunogenic epitope is a B cell epitope or T cell epitope derived from pathogen such as Streptococcus pneumonia, Cryptosporidium parvum, HIV, foot and mouth disease virus, influenza virus, Yersinia pestia, etc. The display of the immunogenic epitope is displayed in the immunogenic loop of HBc, whereas the enhanced stability of self-assembled particles is obtained by the presence of at least one heterologous cysteine residue near the carboxy-terminus of the chimer mol. Methods of making and using the chimers are also disclosed.

L4 ANSWER 5 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 2002:31493 HCAPLUS

DOCUMENT NUMBER: 136:101087

TITLE: Attenuated human-bovine chimeric parainfluenza virus

(PIV) vaccines

INVENTOR(S): Skiadopoulos, Mario H.; Collins, Peter L.; Murphy,

Brian R.; Schmidt, Alexander C.

PATENT ASSIGNEE(S): The Government of the United States of America, as

Represented by the Department of Health and Human

Services, USA

SOURCE: PCT Int. Appl., 154 pp.

CODEN: PIXXD2

Patent

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO. DATE
     PATENT NO.
                         KIND DATE
                                                 -----
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                                -----
                                20020110
                                               WO 2001-US21527 20010705
     WO 2002002605
                         A2
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
              LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
              SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
              YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
               BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                 AU 2001-71909
                                                                     20010705
     AU 2001071909
                          A5
                                20020114
                                20030501
     US 2003082209
                                                 US 2001-900112
                                                                     20010705
                          A1
                                              US 2000-215809P P
                                                                     20000705
PRIORITY APPLN. INFO .:
                                              WO 2001-US21527 W 20010705
```

AB Chimeric human-bovine parainfluenza viruses (PIVs) are infectious and attenuated in humans and other mammals and useful individually or in combination in vaccine formulations for eliciting an anti-PIV immune response. Also provided are isolated polynucleotide mols. and vectors incorporating a chimeric PIV genome or antigenome which includes a partial or complete human or bovine PIV "background" genome or antigenome combined or integrated with one or more heterologous gene(s) or genome segment(s) of a different PIV. Chimeric human-bovine PIV of the invention include a partial or complete "background" PIV genome or antigenome derived from or patterned after a human or bovine PIV virus combined with one or more heterologous gene(s) or genome segment(s) of a different PIV virus to form the human-bovine chimeric PIV genome or antigenome. In certain aspects of the invention, chimeric PIV incorporate a partial or complete human PIV

background genome or antigenome combined with one or more heterologous gene(s) or genome segment(s) from a bovine PIV, whereby the resultant chimeric virus is attenuated by virtue of host-range restriction. In alternate embodiments, human-bovine chimeric PIV incorporate a partial or complete bovine PIV background genome or antigenome combined with one or more heterologous gene(s) or genome segment(s) from a human PIV gene that encode a human PIV immunogenic protein, protein domain or epitope, for example encoded by PIV HN and/or F glycoprotein gene(s) or genome segment(s). Human-bovine chimeric PIV of the invention are also useful as vectors for developing vaccines against other pathogens. A variety of addnl. mutations and nucleotide modifications are provided within the human-bovine chimeric PIV of the invention to yield desired phenotypic and structural effects.

ANSWER 6 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 2001:283263 HCAPLUS

DOCUMENT NUMBER: 135:59874

TITLE:. Chimeric Subgroup A Respiratory

Syncytial Virus with the

Glycoproteins Substituted by Those of Subgroup B and

RSV without the M2-2 Gene Are Attenuated in African Green Monkeys

AUTHOR(S): Cheng, Xing; Zhou, Helen; Tang, Roderick S.; Munoz,

Mary G.; Jin, Hong

CORPORATE SOURCE: Aviron, Mountain View, CA, 94043, USA

SOURCE: Virology (2001), 283(1), 59-68

CODEN: VIRLAX; ISSN: 0042-6822

PUBLISHER: Academic Press

DOCUMENT TYPE: Journal LANGUAGE: English

Using the existing reverse genetics system developed for the subgroup A respiratory syncytial virus (RSV), a chimeric virus (designated rA-GBFB) that expresses subgroup B-specific antigens was constructed by replacing the G and F genes of the A2 strain with those of the 9320 strain of subgroup B RSV. RA-GBFB grew well in tissue culture, but it was attenuated in the respiratory tracts of cotton rats and African green monkeys. To further attenuate this chimeric RSV, the M2-2 open reading frame was removed from rA-GBFB. RA-GBFB.DELTA.M2-2 was highly attenuated in replication in the respiratory tracts of the infected monkeys, but it provided complete protection against wild-type subgroup B RSV challenge following two doses of infection. In this study, rA2.DELTA.M2-2 (a recombinant A2 RSV that lacks the M2-2 gene) was also evaluated in African green monkeys. replication of rA2.DELTA.M2-2 was highly restricted in both the upper and lower respiratory tracts of the infected monkeys and it induced titers of serum anti-RSV neutralizing antibody that were slightly lower than those induced by wild-type rA2. When rA2.DELTA.

M2-2-infected monkeys were challenged with wild-type A2 virus, the replication of the challenge virus was reduced by approx. 100-fold in the upper respiratory tract and 45,000-fold in the lower respiratory tracts.

RA2.DELTA.M2-2 and rA-GBFB.DELTA.M2-2 could represent

a bivalent RSV vaccine compn. for protection against multiple strains from the two RSV subgroups. (c) 2001 Academic Press.

REFERENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 2001:228735 HCAPLUS

DOCUMENT NUMBER: 134:271226

TITLE:

Use of an outer membrane protein A of an enterobacterium associated with a respiratory

syncytial virus immunogenic

peptide for preparing vaccines for intranasal

administration

INVENTOR(S):

Corvaiea, Nathalie; Goestch, Liliane

Pierre Fabre Medicament, Fr. PATENT ASSIGNEE(S):

SOURCE:

PCT Int. Appl., 38 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.		KI	ND	DATE			A:	PPLI	CATI	ои ис	ο.	DATE			
WO 200			A.		2001				20	00-F	R262	6·	2000	0922		
W:	AU, : AT,	BR,	CA,	CN,	JP, DE.	MX, DK.	US, ES.	ZA FI.	FR.	·GB.	GR.	IE,	IT,	LU,	MC,	NL,
144	PT,		J,		•		,									•
FR 279			A.	_	2001			F	R 19	99-1	1888		1999	0923		
BR 200				<del>,</del>	2003			B	R 20	00-1	4246		2000	0922		
EP 121			A.	_	2002					00-9			2000			
R:	AT,	BE, FI.		DΕ,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,

A 19990923 PRIORITY APPLN. INFO.: FR 1999-11888 WO 2000-FR2626 W 20000922

The invention concerns the use of an outer membrane protein A (OmpA) of an enterobacterium, in particular protein P40 of Klebsiella pneumoniae, assocd. with an immunogenic peptide derived from the respiratory syncytial virus (RSV)

for prepg. a pharmaceutical compn. for intranasal administration designed to induce an immune response protecting the upper and lower (lungs) respiratory route against RSV infection. The invention also concerns the use of said compds. for prepg. a vaccine for preventing and treating RSV infection.

REFERENCE COUNT:

THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

11

ACCESSION NUMBER:

2001:50779 HCAPLUS

DOCUMENT NUMBER:

134:114850

TITLE:

Production of recombinant respiratory syncytial viruses expressing immune

modulatory molecules

INVENTOR(S):

Collins, Peter L.; Bukreyev, Alexander; Murphy, Brian

R.; Whitehead, Stephen S.

PATENT ASSIGNEE(S):

United States Dept. of Health and Human Services, USA

PCT Int. Appl., 154 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001004271	A2	20010118	WO 2000-US19042	20000712
WO 2001004271	A3	20010719		

دائعی

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AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS; MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                       Α5
                            20010130
                                            AU 2000-62112
                                                              20000712
     AU 2000062112
     EP 1194581
                       A2
                            20020410
                                            EP 2000-948641
                                                              20000712
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                            20020924
                                            BR 2000-13202
                                                              20000712
     BR 2000013202
                       Α
                       T2
                            20030408
     JP 2003512817
                                            JP 2001-509475
                                                              20000712
PRIORITY APPLN. INFO.:
                                         US 1999-143425P P
                                                              19990713
                                         WO 2000-US19042 W
                                                              20000712
     Recombinant respiratory syncytial virus (
AΒ
    RSV) are provided which express one or more immune modulatory
     mols. The recombinant virus is modified by addn. or substitution of a
     sequences encoding the immune modulatory mol. (e.g., cytokines).
     Introduction of a cytokine increases, decreases, or otherwise enhances
     aspects of viral biol. and/or host immune responses to RSV. In
     one example, the murine interferon-.gamma. gene was inserted into the
     RSV G-F intergenic region. Cultured cells
     infected with rRSV/mIFN-.gamma. expressed the cytokine and replication of
     the recombinant virus was attenuated in upper and lower respiratory tract
     of infected mice.
    ANSWER 9 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN
                         2000:678573 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         133:333659
TITLE:
                         Recombinant respiratory syncytial
                         virus that does not express the NS1
                         or M2-2 protein is highly attenuated and
                         immunogenic in chimpanzees
                         Teng, Michael N.; Whitehead, Stephen S.; Bermingham,
AUTHOR(S):
                         Alison; St. Claire, Marisa; Elkins, William R.;
                         Murphy, Brian R.; Collins, Peter L.
CORPORATE SOURCE:
                         Respiratory Viruses Section, Laboratory of Infectious
                         Diseases, National Institute of Allergy and Infectious
                         Diseases, Bethesda, MD, 20892, USA
SOURCE:
                         Journal of Virology (2000), 74(19), 9317-9321
                         CODEN: JOVIAM; ISSN: 0022-538X
PUBLISHER:
                         American Society for Microbiology
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
AΒ
     Mutant recombinant respiratory syncytial
```

Mutant recombinant respiratory syncytial
viruses (RSV) which cannot express the NS1 and
M2-2 proteins, designated rA2.DELTA.NS1 and rA2.DELTA.
M2-2, resp., were evaluated as live-attenuated RSV
vaccines. The rA2.DELTA.NS1 virus contains a large deletion
that should have the advantageous property of genetic stability during
replication in vitro and in vivo. In vitro, rA2.DELTA.NS1
replicated approx. 10-fold less well than wild-type recombinant
RSV (rA2), while rA2.DELTA.M2-2 had delayed growth
kinetics but reached a final titer similar to that of rA2. Each virus was
administered to the respiratory tracts of RSV-seroneg.
chimpanzees to assess replication, immunogenicity, and
protective efficacy. The rA2.DELTA.NS1 and rA2.DELTA.M2

(IL)

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-2 viruses were 2,200- to 55,000-fold restricted in replication in the
     upper and lower respiratory tracts but induced a level of RSV
     -neutralizing antibody in serum that was only slightly reduced compared to
     the level induced by wild-type RSV. The replication of
     wild-type RSV in immunized chimpanzees after challenge was
     reduced more than 10,000-fold at each site. Importantly, rA2.DELTA.
    NS1 and rA2.DELTA.M2-2 were 10-fold more restricted in
     replication in the upper respiratory tract than was the cpts248/404 virus,
     a vaccine candidate that retained mild reactogenicity in the upper
     respiratory tracts of 1-mo-old infants. Thus, either rA2.DELTA.
    NS1 or rA2.DELTA.M2-2 might be appropriately attenuated
     for this age group, which is the major target population for an
     RSV vaccine. In addn., these results show that neither
     NS1 nor M2-2 is essential for RSV replication
     in vivo, although each is important for efficient replication.
                               THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                         34
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 10 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN
                         1997:775451 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         128:60477
                         Recombinant vaccinia viruses expressing the F
TITLE:
                         , G or N, but not the M2, protein
                         of bovine respiratory syncytial
                         virus (BRSV) induce resistance to BRSV
                         challenge in the calf and protect against the
                         development of pneumonic lesions
                         Taylor, Geraldine; Thomas, Lewis H.; Furze, Julie M.;
AUTHOR(S):
                         Cook, Roy S.; Wyld, Sara G.; Lerch, Robert; Hardy,
                         Richard; Wertz, Gail W.
                        Institute for Animal Health, Newbury, RG20 7NN, UK
CORPORATE SOURCE:
                         Journal of General Virology (1997), 78(12), 3195-3206
SOURCE:
                         CODEN: JGVIAY; ISSN: 0022-1317
PUBLISHER:
                         Society for General Microbiology
                         Journal
DOCUMENT TYPE:
LANGUAGE:
                         English
     The immunogenicity and protective efficacy of recombinant
     vaccinia viruses (rVV) encoding the F, G, N, or
    M2 (22K) proteins of bovine respiratory
     syncytial virus (BRSV) were evaluated in calves, the
     natural host for BRSV. Calves were vaccinated either by scarification or
     intratracheally with rVV and challenged 6-7 wk later with BRSV. Although
     replication of rVV expressing the F protein in the respiratory
     tract was limited after intratracheal vaccination, the levels of serum and
     pulmonary antibody were similar to those induced following scarification.
     The serum antibody response induced by the F protein was biased
     in favor of IgG1 antibody, whereas the G and the N proteins
     induced similar levels of IgG1: IgG2, and antibody was undetectable in
     calves primed with the M2 protein. The F protein
     induced neutralizing antibodies, but only levels of complement-dependent
     neutralizing antibodies were induced by the G protein, and
     antibody induced by the N protein was not neutralizing. The F
```

and N proteins primed calves for BRSV-specific lymphocyte proliferative responses, whereas proliferative responses were detected in calves primed

against BRSV infection and, in contrast with the enhanced lung pathol. seen in mice vaccinated with rVV expressing individual proteins of human

differences in the immune responses induced by the rVVs, the F,

protein primed lymphocytes in only 1 out of 5 calves. Although there were

with the G protein only after BRSV challenge. The M2

G and N, but not the M2, proteins induced protection

RSV, there was a redn. in lung pathol. in calves.

35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS . REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 11 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

1997:744681 HCAPLUS ACCESSION NUMBER:

128:45651 DOCUMENT NUMBER:

TITLE:

(A)

Recombinant respiratory syncytial virus from which the entire SH gene

has been deleted grows efficiently in cell culture and exhibits site-specific attenuation in the respiratory

tract of the mouse

AUTHOR(S):

Bukreyev, Alexander; Whitehead, Stephen S.; Murphy,

Brian R.; Collins, Peter L.

Laboratory of Infectious Diseases, National Institute CORPORATE SOURCE:

Allergy and Infectious Diseases, Bethesda, MD,

20892-0720, USA

SOURCE:

PUBLISHER:

Journal of Virology (1997), 71(12), 8973-8982

CODEN: JOVIAM; ISSN: 0022-538X American Society for Microbiology

DOCUMENT TYPE:

Journal English LANGUAGE:

The small hydrophobic protein SH of human respiratory ΑB syncytial virus (RSV) is a short transmembrane surface protein of unknown function. A full-length cDNA of RSV strain A2 (subgroup A) antigenomic RNA was modified such that the entire SH gene, including the transcription signals and the complete mRNA-encoding sequence, was deleted and replaced by a synthetic intergenic region. This reduced the length of the antigenome by 398 nucleotides and ablated expression of 1 of the 10 RSV mRNAs. Recombinant virus contg. this engineered deletion was recovered, and the absence of the SH gene was confirmed by reverse transcription in conjunction with Northern blot anal. of intracellular RNAs and gel electrophoresis of labeled intracellular proteins confirmed the lack of expression of the SH mRNA and protein. The absence of the SH gene did not noticeably affect RNA replication, but two effects on transcription were noted. First, synthesis of the G, F, and M2 mRNAs was increased, presumably due to their being one position closer to the promoter in the gene order. Second, transcription of genes downstream of the engineered site exhibited a steeper gradient of polarity. On monolayers of HEp-2 cells, the SH-minus virus produced syncytia which were at least equiv. in size to those of the wild type and produced plaques which were 70% larger. Furthermore, the SH-minus virus grew somewhat better (up to 12.6-fold) than wild-type recombinant RSV in certain cell lines. While the function of the SH protein remains to be detd., it seems to be completely dispensable for growth in tissue culture and fusion function. When inoculated intranasally into mice, the SH-minus virus resembled the wild-type recombinant virus in its efficiency of replication in the lungs, whereas it replicated 10-fold less efficiently in the upper respiratory tract. In mice, the SH-minus and wild-type recombinant viruses were similarly immunogenic and effective in inducing resistance to virus challenge.

REFERENCE COUNT:

THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS 41 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 12 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

1997:739237 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 128:33496

Structural properties of chimeric peptides containing TITLE:

a T-cell epitope linked to a fusion peptide and their importance for in vivo induction of cytotoxic T-cell

responses

AUTHOR(S): Lelievre, Dominique; Hsu, Shiou-Chih; Daubos,

Philippe; Favard, Cyril; Vigny, Paul; Trudelle, Yves;

Steward, Michael W.; Delmas, Agnes

CORPORATE SOURCE: Centre de Biophysique Moleculaire, UPR 4301 CNRS,

Orleans, F-45071, Fr.

SOURCE: European Journal of Biochemistry (1997), 249(3),

895-904

CODEN: EJBCAI; ISSN: 0014-2956

PUBLISHER: Springer
DOCUMENT TYPE: Journal
LANGUAGE: English

AB The authors have previously shown that when administered to mice without adjuvant, a chimeric peptide consisting of the fusion peptide F from measles virus protein linked at the C-terminus of a cytotoxic T-cell epitope from the M2 protein of respiratory

syncytial virus efficiently primes for an major

histocompatibility complex (MHC) class-I restricted cytotoxic T lymphocyte (CTL) response. Here, the authors demonstrated by microspectrofluorometry that the fusion-peptide moiety bound to the plasma membrane of living cells. When the fusion peptide was linked to the C-terminus of the CTL epitope, the chimeric peptide (M2-F) adopted a marked .beta.-sheet conformation. In contrast, when the fusion peptide was linked to the N-terminus of the T-cell epitope (F-M2), the chimeric peptide adopted an .alpha.-helical conformation in the presence of trifluoroethanol. The immunogenicity of the 2 chimeric peptides for class-I restricted CTL was also different, the one adopting the .alpha.-helical conformation being more immunogenic . Probably due to its obvious conversion to an .alpha.-helical conformation, the F-M2 peptide could have a higher

propensity to insert into membranes, as shown by microspectrofluorometry,

.

L4 ANSWER 13 OF 13 HCAPLUS COPYRIGHT 2003 ACS on STN

with a resultant better immunogenicity than the M2-

ACCESSION NUMBER:

F peptide.

1994:6778 HCAPLUS

DOCUMENT NUMBER:

120:6778

TITLE:

CAS

Mutant respiratory syncytial

virus (RSV), vaccines containing it,
and methods of vaccination with RSV

INVENTOR(S):

Randolph, Valerie Bruce; Crowley, Joan Coflan

PATENT ASSIGNEE(S): American Cyanamid Co., USA SOURCE: Eur. Pat. Appl., 63 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 567100	A1	19931027	EP 1993-106496	19930421
EP 567100	B1	19990317	B1 1995 100190	15550121
R: AT, BE,	CH, DE	, DK, ES, FR,	GB, GR, IE, IT, LI	, LU, NL, PT, SE
CA 2094464	AA	19931022	CA 1993-2094464	19930420
ZA 9302763	A	19940208	ZA 1993-2763	19930420
AU 9337057	A1	19931028	AU 1993-37057	19930421
AU 671983	B2	19960919		

19940201 19930421 JP 06022756 A2 JP 1993-117812 AT 177785 19990415 AT 1993-106496 19930421 E ES 2130189 Т3 19990701 ES 1993-106496 19930421 US 1992-871420 19920421 PRIORITY APPLN. INFO.: AB Cold-adapted (attenuated) mutants of RSV belonging to subgroup A or B were obtained by repeated passage in Vero cells at .ltoreq.26.degree... The immunogenic peptides are purified and identified, and the nucleic acid segments encoding them are cloned, sequenced, and expressed. The mutants elicited protective immunity against RSV challenge in cotton rats and African green monkeys. Monoclonal antibodies to the mutants are useful in diagnostic assays and therapy.

#### => d ibib abs 16 1-19

L6 ANSWER 1 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER: 2003-559095 [52] WPIDS

DOC. NO. CPI: C2003-150701

TITLE:

a

An immunogenic composition comprising an antigen and a purified M protein from

respiratory syncytial virus,

useful as a vaccine for immunizing a host or for

enhancing the immune response to an antigen in a host.

DERWENT CLASS: B04 D16

INVENTOR(S): BARBER, B; CATES, G; PARRINGTON, M; SAMBHARA, S

PATENT ASSIGNEE(S): (AVET) AVENTIS PASTEUR LTD

COUNTRY COUNT: 102

PATENT INFORMATION:

PATENT NO	KIND	DATE	WEEK	LA	PG

WO 2003053464 A1 20030703 (200352)\* EN 27

RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SI SK SL SZ TR TZ UG ZM ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR GU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ OM PH PL PT RO RU SC SD SE SG SK SL TJ TM TN TR TT TZ UA UG US UZ VC VN YU ZA ZM ZW

#### APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION DATE	
WO 20030534	64 A1	WO 2002-CA1953 20021218	

PRIORITY APPLN. INFO: US 2001-341422P 20011220

AN 2003-559095 [52] WPIDS

AB WO2003053464 A UPAB: 20030813

NOVELTY - An immunogenic composition comprising an antigen and a purified M protein from respiratory syncytial . virus or at least one of its immunoeffective fragments, is new. The M protein or its immunoeffective fragment is provided in a pre-selected amount to provide an enhanced immune response to the antigen

in a host having a pre-existing respiratory syncytial virus M-specific immune response.

 ${\tt DETAILED}$  <code>DESCRIPTION</code> - <code>INDEPENDENT</code> <code>CLAIMS</code> are included for the following:

(1) making an immunogenic composition comprising providing an antigen and a purified M protein from respiratory

syncytial virus or at least one of its immunoeffective
fragment, where the amount of the M protein or its
immunoeffective fragment is provided in a pre-selected amount to provide
an enhanced immune response to the antigen in a host having a pre-existing
respiratory syncytial virus M
-specific immune response;

- (2) immunizing a host comprising administering the immunogenic composition cited above in a host; and
- (3) enhancing an immune response to an antigen in a host having a pre-existing immune response to respiratory syncytial virus M protein comprising purifying M protein of respiratory syncytial virus, mixing a pre-selected amount of the purified M protein with a different antigen, formulating the mixture as a vaccine, and administering the vaccine to a host.

ACTIVITY - Virucide.

MECHANISM OF ACTION - Vaccine.

Groups of BALB/c mice were primed with 103 pfu of live respiratory syncytial virus (RSV)

intranasally. Four weeks later, the mice were boosted with phosphate buffered saline, or formalin-inactivated-RSV 5 ng of F with 1 micro g of M in aluminum phosphate or 5 ng of F in aluminum phosphate. The animals were boosted again four weeks later and the sera samples were collected to determine anti-F responses by F-specific enzyme-linked immunosorbent assay. The antibody results show that the inclusion of the RSV M antigen enhances the antibody response (antibody titer) to RSV F antigen by two logs when compared to the RSV F only boost.

USE - The immunogenic composition is useful as a vaccine for immunizing a host against a disease caused by respiratory syncytial virus. The pre-selected amount of purified M protein from respiratory syncytial virus or its immunoeffective fragment is useful for enhancing the immune response to an antigen in a host having a pre-existing respiratory syncytial virus M -specific immune response (claimed).

Dwg.0/4

.6 ANSWER 2 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2003-381589 [36] WPIDS

DOC. NO. CPI:

C2003-101341

TITLE:

New immunogenic compositions comprising a cocktail of at least four different RSV

antigens, useful as a vaccine against respiratory

syncytial virus (RSV), which

causes respiratory tract infections in infants and

children.

DERWENT CLASS:

B04 D16

INVENTOR(S):

HUANG, S; KUMAR, M; LEONG, K; MOHAPATRA, S S; BEHERA, A K; CHEN, L; LEONG, K W; LOCKEY, R F; MOHAPTRA, S S; PEREZ

DE LA CRUZ, C; ZHANG, J

PATENT ASSIGNEE(S):

(HUAN-I) HUANG S; (KUMA-I) KUMAR M; (LEON-I) LEONG K; (MOHA-I) MOHAPATRA S S; (UYJO) UNIV JOHNS HOPKINS;

(UYSF-N) UNIV SOUTH FLORIDA

COUNTRY COUNT:

9'4

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2003028759 A1 20030410 (200336)\* EN 35

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

NL OA PT SD SE SL SZ TR TZ UG ZM ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK

LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG

SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

US 2003068333 A1 20030410 (200340)

#### APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 2003028 US 2003068	759 Al 333 Al Provisional	WO 2002-US4114 US 2001-325573P US 2002-73065	20020212 20010928 20020212

PRIORITY APPLN. INFO: US 2001-325573P 20010928; US 2002-73065

20020212

AN 2003-381589 [36] WPIDS

AB W02003028759 A UPAB: 20030609

NOVELTY - Immunogenic compositions for conferring protection in

a host against disease caused by respiratory syncytial

virus (RSV) comprising an F RSV

antigen and a G RSV antigen, or an M2

RSV antigen; or an F RSV antigen, a G

RSV antigen and an M2 RSV antigen, and at

least one of M, M2, SH, NS1,

NS2, N, F, G, or P RSV

antigen, are new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) a gene expression vaccine for conferring protection in a host against disease caused by RSV comprising a plasmid DNA cocktail comprising a combination of at least two RSV antigens selected from F, G, M, M2, SH,

NS1, NS2, N, and P; where the plasmid DNA

cocktail is coacervated with chitosan to form nanospheres;

(2) immunizing a host against disease caused by infection with RSV; and

(3) making a gene expression vaccine.

ACTIVITY - Virucide.

BALB/c mice were orally administered with the composition or naked DNA (25 micro g total). Animals were infected with RSV on day 16, and 4 days later sacrificed. Results showed that mice given the composition had reduction in epithelial cell damage and interstitial thickening when compared to controls.

MECHANISM OF ACTION - Vaccine.

USE - The composition is useful as a vaccine against respiratory syncytial virus (RSV), which causes respiratory tract infections in infants and children. Dwg.0/9

L6 ANSWER 3 OF 19 MEDLINE on STN

ACCESSION NUMBER: 2003386614 'IN-PROCESS

DOCUMENT NUMBER: 22804932 PubMed ID: 12922094

TITLE: Evaluation of recombinant respiratory syncytial virus gene deletion mutants in

African green monkeys for their potential as live

Lucas 10/073,065 25/08/2003

attenuated vaccine candidates.

Jin Hong; Cheng Xing; Traina-Dorge Vicki L; Park Hyun Jung; AUTHOR:

Zhou Helen; Soike Ken; Kemble George

MedImmune Vaccines Inc., 297 North Bernardo Avenue, 94043, CORPORATE SOURCE:

Mountain View, CA, USA.

VACCINE, (2003 Sep 8) 21 (25-26) 3647-52. SOURCE:

Journal code: 8406899. ISSN: 0264-410X.

Netherlands PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

English LANGUAGE:

IN-PROCESS; NONINDEXED; Priority Journals FILE SEGMENT:

Entered STN: 20030819 ENTRY DATE:

Last Updated on STN: 20030819

Towards the goal of developing live attenuated respiratory AB syncytial virus (RSV) vaccines to prevent

severe respiratory tract infections caused by respiratory

syncytial virus, recombinant RSV containing a deletion of single or multiple NS1, NS2, SH

and M2-2 genes have been generated. In this study,

recombinants, rA2DeltaM2-2, rA2DeltaNS2, rA2DeltaNS1NS2, rA2DeltaSHNS2, rA2DeltaM2-2NS2 were evaluated in African green monkeys (AGMs) for their

infectivity, immunogenicity and protection against wild type

(wt) RSV challenge. Replication of rA2DeltaNS2 and

rA2DeltaSHNS2 was not attenuated in either the upper or the lower respiratory tracts of AGMs. On the other hands, rA2DeltaNS1NS2 was over-attenuated; it did not replicate in the respiratory tracts of the infected monkeys and did not provide sufficient protection against wild type RSV challenge. rA2DeltaM2-2NS2 was slightly more attenuated than rA2DeltaM2-2 and provided partial protection against wt RSV

challenge. rA2DeltaM2-2, and possibly rA2DeltaM2-2NS2, exhibited the attenuated but protective phenotypes in the monkeys that could be further evaluated as potential live attenuated RSV vaccine candidates in

the clinical studies.

ANSWER 4 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

2002-090518 [12] ACCESSION NUMBER:

DOC. NO. CPI: C2002-027998

TITLE: An isolated infectious recombinant respiratory

syncytial virus (RSV) having

one or more shifted RSV gene(s) or genome

WPIDS

segment(s) within the recombinant genome or antigenome,

useful as an attenuated vaccine against RSV

strains.

DERWENT CLASS: B04 D16

BUCHHOLZ, U; COLLINS, P L; KREMPL, C D; MURPHY, B R; INVENTOR(S):

WHITEHEAD; S S

(USGO) US GOVERNMENT; (BUCH-I) BUCHHOLZ U; (COLL-I) PATENT ASSIGNEE(S):

COLLINS P L; (KREM-I) KREMPL C D; (MURP-I) MURPHY B R;

(WHIT-I) WHITEHEAD S S

COUNTRY COUNT: 96

PATENT INFORMATION:

WEEK LA PATENT NO KIND DATE \_\_\_\_\_\_

WO 2002000693 A2 20020103 (200212)\* EN 168

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK

DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ

LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD

SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001068709 A 20020108 (200235)

US 2002146433 A1 20021010 (200269)

EP 1294858 A2 20030326 (200323) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

#### APPLICATION DETAILS:

PATENT NO K	IND	APPLICATION	DATE
WO 2002000693 AU 2001068709 US 2002146433 EP 1294858		WO 2001-US20107 AU 2001-68709 US 2000-213708P US 2001-887469 EP 2001-946696 WO 2001-US20107	20010622 20010622 20000623 20010622 20010622 20010622
		MO 5001-0250101	20010022

#### FILING DETAILS:

PAT	TENT NO	KIND			PAT	ENT NO
ΑU	2001068709	9 A	Based	on	WO	200200693
ΕP	1294858	A2	Based	on	WO	200200693

PRIORITY APPLN. INFO: US 2000-213708P 20000623; US 2001-887469 20010622

AN 2002-090518 [12] WPIDS

AB WO 200200693 A UPAB: 20020221

NOVELTY - An isolated infectious recombinant respiratory syncytial virus (RSV) having one or more shifted RSV gene(s) or genome segment(s) within the recombinant genome or antigenome that is/are positionally shifted to a more promoter-proximal or promoter-distal position relative to a position of the RSV gene(s) or genome segment(s) within a wild type RSV genome or antigenome, is new.

DETAILED DESCRIPTION - An isolated infectious recombinant respiratory syncytial virus (RSV) comprising a major nucleocapsid (N) protein, a nucleocapsid phosphoprotein (P), a large polymerase protein (L), a RNA polymerase elongation factor, and a partial or complete recombinant RSV genome or antigenome having one or more shifted RSV gene(s) or genome segment(s) within the recombinant genome or antigenome that is/are positionally shifted to a more promoter-proximal or promoter-distal position relative to a position of the RSV gene(s) or genome segment(s) within a wild type RSV genome or antigenome.

INDEPENDENT CLAIMS are included for the following:

- (1) a method (M1) for stimulating the immune system of an individual to induce protection against RSV which comprises administering to the individual an immunologically sufficient amount of the recombinant RSV combined with a physiologically acceptable carrier;
- (2) an isolated polynucleotide molecule comprising a recombinant RSV genome or antigenome having one or more shifted RSV gene(s) or genome segment(s) within the recombinant genome or antigenome that is/are positionally shifted to a more promoter-proximal or promoter-distal position relative to a position of the RSV gene(s) or genome segment(s) within a wild type RSV genome or antigenome;
- (3) a method (M2) for producing an infectious attenuated recombinant RSV particle from one or more isolated

polynucleotide molecules encoding the RSV, comprising expressing in a cell or cell-free lysate an expression vector comprising an isolated polynucleotide comprising a recombinant RSV genome or antigenome having one or more shifted RSV gene(s) or genome segment(s) within the recombinant genome or antigenome that is/are positionally shifted to a more promoter-proximal or promoter-5 distal position relative to a position of the RSV gene(s) or genome segment(s) within a wild type RSV genome or antigenome, and RSV N, P, L and RNA polymerase elongation factor proteins;

- (4) an isolated infectious chimeric RSV comprising a major nucleocapsid (N) protein, a nucleocapsid phosphoprotein (P), a large polymerase protein (L), a RNA polymerase elongation factor, and a partial or complete bovine RSV background genome or antigenome combined with heterologous gene(s) and/or genome segment(s) of a human RSV selected from heterologous gene(s) and/or genome segment(s) of RSV NS1, NS2, M, SH,
- G, and/or F, to form a human-bovine chimeric RSV
  genome or antigenome; and
- (5) an isolated polynucleotide molecule comprising a recombinant RSV genome or antigenome comprising a partial or complete bovine RSV background genome or antigenome combined with a plurality of heterologous gene(s) and/or genome segment(s) of a human RSV selected from heterologous gene(s) and/or genome segment(s) of RSV NS1, NS2, M, SH, G, and/or

F genes, to form a human-bovine chimeric RSV genome or antigenome.

ACTIVITY - Antiviral.

No biological data given.

MECHANISM OF ACTION - The recombinant RSV elicits an immune response against either human RSV A or RSV B or both human RSV A and RSV B (claimed); gene therapy; vaccine.

No biological data given.

USE - The recombinant RSV is useful in an attenuated vaccine to elicits an immune response against one or more strains of RSV.

Dwg.0/15

L6 ANSWER 5 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2003-128101 [12] WPIDS...

CROSS REFERENCE:

1998-110527 [10]

DOC. NO. NON-CPI:

N2003-101703

DOC. NO. CPI:

C2003-032723

TITLE:

Composition for vaccination, diagnosis and treatment of

respiratory syncytial virus

infection, comprises fusion protein, attachment protein

and matrix protein of respiratory

syncytial virus.

DERWENT CLASS:

A96 B04 D16 S03

INVENTOR(S): CATES, G A; KLI

CATES, G A; KLEIN, M H; OOMEN, R P; SANHUEZA, S E

(CATE-I) CATES G A; (KLEI-I) KLEIN M H; (OOME-I) OOMEN R P; (SANH-I) SANHUEZA S E; (AVET) AVENTIS PASTEUR LTD

101

COUNTRY COUNT:

PATENT ASSIGNEE(S):

PATENT INFORMATION:

PATENT N	NO KI	ND DATE	WEEK	LA	PG

US 2002136739 A1 20020926 (200312)\* 23 WO 2003022878 A2 20030320 (200330) EN

RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GH GM GR IE IT KE LS LU

MC MW MZ NL OA PT SD SE SK SL SZ TR TZ UG ZM ZW
W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK
DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR
KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ OM PH PL PT
RO RU SD SE SG SI SK SL TJ TM TN TR TT TZ UA UG US UZ VC VN YU ZA
ZM ZW

#### APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
US 2002136739 A1 CIP of CIP of CIP of	US 1996-679060 WO 1997-CA497 US 1999-214605	19960712 19970711 19990503
WO 2003022878 A2	US 2001-950655 WO 2002-CA1347	20010913 20020903

#### FILING DETAILS:

PATENT NO	KIND	PATENT NO
	739 A1 CIP of	US 6020182 US 6309649

PRIORITY APPLN. INFO: US 2001-950655 20010913; US 1996-679060 19960712; WO 1997-CA497 19970711; US 1999-214605 19990503

AN 2003-128101 [12] WPIDS

CR 1998-110527 [10]

AB US2002136739 A UPAB: 20030513

NOVELTY - A mixture (I) of purified fusion (F) protein, attachment (G) protein and matrix (M) protein of respiratory syncytial virus (RSV), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for:

- (1) an immunogenic composition (II) comprising (I);
- (2) producing (M1) a vaccine for protection against RSV, by administering (II) to a test host to determine the amount of frequency of administration to confer protection against disease caused by RSV , and formulating the immunogenic composition in a form suitable for administration to a treated host in accordance with the determined amount and frequency of administration;
- (3) producing (M2) a coisolated and copurified mixture of proteins of RSV (III), by growing RSV on cells in a culture medium, separating the grown virus from the culture medium, solubilizing (F), (G) and (M) protein from the separated virus, and coisolating and copurifying the solubilized RSV proteins; and
- (4) a diagnostic kit (IV) for determining the presence of antibodies in a sample specifically reactive with F, G or M protein of RSV, comprising (I), unit for contacting the immunogenic composition with the sample to produce complexes comprising RSV and any antibodies present in the sample and unit for determining the production of the complexes.

ACTIVITY - Virucide.

MECHANISM OF ACTION - Vaccine.

RSV subunit preparations were used to formulate an alum-adjuvanted vaccine and a placebo control that contained only alum. The total protein present in a single dose of the vaccines of the antigens RSV  $\mathbf{F}$ ,  $\mathbf{G}$  and  $\mathbf{M}$  was 100 micro

g, present in 0.5 ml of phosphate buffered saline. In the alum-adjuvanted vaccine, there was 1.5 mg of alum/0.5 ml of vaccine. The vaccines were assessed for stability for 42 months at 5 deg. C, 5 months at 25 deg. C and 5 weeks at 37 deg. C to ensure physical and biological stability over time. Stability studies indicated that the P and G antigens in the alum-adjuvanted vaccines were stable at 25 deg. C for at least 6 weeks. The vaccine preparations were used to immunize adults, 65 years of age or older. Blood samples were obtained on day 0 (day of immunization), day 32, day 60 and day 180 and RSV serology was performed on the serum samples. RSV neutralization assay was performed by a plaque reduction method (NA) against RSV A and RSV B. There was a greater or equal to 2-fold increase in antibody titer or 4-fold increase in antibody titer compared to pre-immunization titers.

USE - (I) is useful as a pharmaceutical substance in a vaccine against disease caused by infection with respiratory syncytial virus. (I) and (II) are useful for determining the presence in a sample of antibodies specifically reactive with F, G or M protein of RSV, by contacting the sample with (I) to produce complexes comprising RSV and any antibodies present in the sample specifically reactive with it, or immunizing a subject with (II) to produce antibodies specific for the proteins and contacting the sample with antibodies to produce complexes, and determining the production of the complexes. (II) is formulated as a vaccine for in vivo administration to a host, especially a primate, human to confer protection against RSV. (II) is useful for generating an immune response in a host. (II) is useful for producing monoclonal antibodies specific for (F), (G) and (M) protein of RSV, by administering the immunogenic composition to at least one mouse to produce at least one immunized mouse, removing B-lymphocytes from the immunized mouse, fusing the B-lymphocytes with myeloma cells, producing hybridomas, cloning the hybridomas which produce a selected anti-RSV protein antibody, culturing the selected anti-RSV protein antibody-producing clones, and isolating anti-RSV protein antibodies from the selected cultures (all claimed).

ADVANTAGE - (I) as a vaccine is safe and highly immunogenic

Dwg.0/6

L6 ANSWER 6 OF 19 MEDLINE on STN DUPLICATE 1

ACCESSION NUMBER: 2002654767 MEDLINE

DOCUMENT NUMBER: 22302303 PubMed ID: 12414935
TITLE: Recombinant respiratory syncytial

virus with the G and F genes

shifted to the promoter-proximal positions.

AUTHOR: Krempl Christine; Murphy Brian R; Collins Peter L

CORPORATE SOURCE: Laboratory of Infectious Diseases, National Institute of

Allergy and Infectious Diseases, Bethesda, Maryland

20892-8007, USA.

CONTRACT NUMBER: AI-00087 (NIAID)

SOURCE: JOURNAL OF VIROLOGY, (2002 Dec) 76 (23) 11931-42.

Journal code: 0113724. ISSN: 0022-538X.

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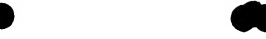
LANGUAGE: English

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ENTRY MONTH: 200212

ENTRY DATE: Entered STN: 20021105

Last Updated on STN: 20021218



Entered Medline: 20021213

The genome of human respiratory syncytial AB virus (RSV) encodes 10 mRNAs and 11 proteins in the order 3'-NS1-NS2-N-P-M-SH -G-F-M2-1/M2-2-L-5'. The G and F glycoproteins are the major RSV neutralization and protective antigens. It seems likely that a high level of expression of G and F would be desirable for a live RSV vaccine. For mononegaviruses, the gene order is a major factor controlling the level of mRNA and protein expression due to the polar gradient of sequential transcription. In order to increase the expression of G and F, recombinant RSVs based on strain A2 were constructed in which the G or F gene was shifted from the sixth or seventh position (in a genome lacking the SH gene), respectively, to the first position (rRSV-G1/DeltaSH and rRSV-F1/DeltaSH, respectively). Another virus was made in which  ${\bf G}$  and  ${\bf F}$  were shifted together to the first and second positions, respectively (rRSV-G1F2/DeltaSH). Shifting one or two genes to the promoter-proximal position resulted in increased mRNA and protein expression of the shifted genes, with G and F expression increased up to 2.4-and 7.8-fold, respectively, at the mRNA level and approximately 2.5-fold at the protein level, compared to the parental virus. Interestingly, the transcription of downstream genes was not greatly affected even though shifting G or F, or G and F together, had the consequence of moving the block of genes NS1-NS2-N-P-M-(G) one or two positions further from the promoter. The efficiency of replication of the gene shift viruses in vitro was increased up to 10-fold. However, their efficiency of replication in the lower respiratory tracts of mice was statistically indistinguishable from that of the parental virus. In the upper respiratory tract, replication was slightly reduced on some days for viruses in which G was in

the first position. The magnitude of the **G**-specific antibody response to the gene shift viruses was similar to that to the parental virus, whereas the **F**-specific response was increased up to fourfold, although this was not reflected in an increase of the neutralizing activity. Thus, shifting the **G** and **F** genes to the promoter-proximal position increased virus replication in vitro, had little effect on replication in the mouse, and increased the antigen-specific **immunogenicity** of the virus beyond that of parental **RSV**.

L6 ANSWER 7 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2001-356173 [37] WPIDS

CROSS REFERENCE:

1999-045317 [04]; 2001-091890 [10]

DOC. NO. CPI:

C2001-110518

TITLE:

Isolated infectious chimeric parainfluenza virus (PIV), useful in an attenuated vaccine to elicits an immune response against one or more virus(es) selected from

human PIV1 (HPIV1), HPIV2 and HPIV3.

DERWENT CLASS:

B04 D16 P32

INVENTOR(S):

COLLINS, P L; DURBIN, A P; MURPHY, B R; SCHMIDT, A C;

SKIADOPOULOS, M H; TAO, T

PATENT ASSIGNEE(S):

(USSH) US DEPT HEALTH & HUMAN SERVICES; (COLL-I) COLLINS P L; (DURB-I) DURBIN A P; (MURP-I) MURPHY B R; (SCHM-I) SCHMIDT A C; (SKIA-I) SKIADOPOULOS M H; (TAOT-I) TAO T

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

95



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WO 2001042445 A2 20010614 (200137)* EN 305
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SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001020731 A 20010618 (200161) A2 20020213 (200219) EN EP 1179054

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL RO SI

A 20020501 (200252) CN 1347453 US 2002155581 A1 20021024 (200273)

JP 2003516148 W 20030513 (200334) 367

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#### APPLICATION DETAILS:

PATENT NO KIND	AP	PLICATION	DATE
WO 2001042445 A2 AU 2001020731 A EP 1179054 A2	AU EP	2000-US33293 2001-20731 2000-984052 2000-US33293	20001208 20001208 20001208 20001208
•	Provisional US Provisional US CIP of US Provisional US	2000-805939 1997-47575P 1997-59385P 1998-83793 1999-170195P	20001208 19970523 19970919 19980522 19991210
JP 2003516148 W	WO	2000-733692 2000-US33293 2001-544321	20001208 20001208 20001208

#### FILING DETAILS:

PAT	TENT NO	KIND			PAT	TENT NO
AU	20010207	31 A	Based	on	WO	200142445
ΕP	1179054	A2	Based	on	WO	200142445
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PRIORITY APPLN. INFO: US 1999-459062 19991210; US 1999-170195P 19991210; US 1999-458813 19991210; US 1997-47575P 19970523; US 1997-59385P 19980522; US 19970919; US 1998-83793 2000-733692 20001208

WPIDS 2001-356173 [37] AN

1999-045317 [04]; 2001-091890 [10] CR

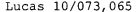
WO 200142445 A UPAB: 20030529 AΒ

NOVELTY - An isolated infectious chimeric parainfluenza virus (PIV), is new.

DETAILED DESCRIPTION - An isolated infectious chimeric parainfluenza virus (PIV), is new.

The virus comprises a major nucleocapsid protein (N), a nucleocapsid phosphoprotein (P), a large polymerase protein (L), and a partial or complete PIV vector.background genome, or antigenome combined with one or more heterologous gene(s) or genome segment(s) encoding one or more antigenic determinants of one or more heterologous pathogen(s) to form a chimeric genome or antigenome.

INDEPENDENT CLAIMS are also included for the following:





- (1) a method for stimulating the immune system of an individual to induce protection against PIV, comprising administering an immunologically sufficient amount of the chimeric PIV;
- (2) a method for sequential immunization to stimulate the immune system of an individual to induce protection against multiple pathogens comprising administering to a newborn to 4 month old infant an immunologically sufficient amount of a first attenuated chimeric human PIV (HPIV) expressing an antigenic determinant of a non-PIV pathogen and one or more antigenic determinants of HPIV3 and subsequently administering an immunologically sufficient amount of a second attenuated chimeric HPIV expressing an antigenic determinant of a non-PIV pathogen and one or more antigenic determinants of HPIV1 or HPIV2;
- (3) an isolated polynucleotide comprising a chimeric PIV genome or antigenome which includes a partial or complete PIV background genome or antigenome combined with one or more heterologous gene(s) or genome segment(s) encoding one or more antigenic determinant(s) of one or more heterologous pathogen(s) to form a chimeric PIV genome or antigenome;
- (4) a method for producing an infectious attenuated chimeric PIV particle from one or more isolated polynucleotide molecules encoding the PIV, comprising expressing in a cell or cell-free lysate an expression vector comprising an isolated polynucleotide comprising a partial or complete PIV vector genome or antigenome of a human or bovine PIV combined with one or more heterologous gene(s) or genome segment(s) encoding one or more antigenic determinant(s) of one or more heterologous pathogen(s) to form a chimeric PIV genome or antigenome, and PIV N, P and L proteins;
- (5) an expression vector comprising an operably linked transcriptional promoter, a polynucleotide sequence which includes a partial or complete PIV vector genome or antigenome of a human or bovine PIV combined with one or more heterologous gene(s) or genome segment(s) encoding one or more antigenic determinant(s) of one or more heterologous pathogen(s) to form a chimeric PIV genome or antigenome, and a transcriptional terminator; and
- (6) an isolated infectious recombinant PIV comprising a N protein, a P, a L, and a PIV genome or antigenome having a polynucleotide insertion of between 150 nucleotides and 4000 nucleotides in length in a non-coding region (NCR) of the genome or antigenome or as a separate gene unit (GU), the polynucleotide insertion lacking a complete open reading frame (ORF) and specifying an attenuated phenotype in the recombinant PIV. ACTIVITY - Antiviral.

Chimpanzees in groups of 4 were inoculated intranasally and intratracheally with 105 TCID50 of rPIV3-2TM or PIV2/V94 on day 0. NT swab specimens (day 1 to 12) and tracheal lavage (days 2, 4, 6, 8, and 10) samples were collected. Virus titer was determined as previously described (Durbin et al., Virology 261:319-30, 1999), and results are expressed as log10 TCID50/ml. rPIV3-2TM had a lower peak titer than its wild type parent PIV2/V94 and was shed for a significantly shorter duration than PIV2/94, indicating that rPIV3-2TM is attenuated in chimpanzees. PIV2/94 wild-type virus replicates to low levels in chimpanzees compared to hamsters and AFGs (undefined), while rPIV3-2TM virus was attenuated in each of these model hosts.

MECHANISM OF ACTION - Anti-PIV vaccine.

USE - The chimeric PIV is useful in an attenuated vaccine to elicits an immune response against one or more virus(es) selected from HPIV1, HPIV2 and HPIV3. Preferably, the chimeric PIV elicits an immune response against HPIV3 and another virus selected from HPIV1 or HPIV2. The chimeric PIV may also elicits a polyspecific immune response against HPIV3 and measles or respiratory syncytial virus. An immunospecific composition may also contain two chimeric PIVs, where the first chimeric PIV elicits an immune response against HPIV3 and the second



chimeric PIV elicits an immune response against HPIV1 or HPIV2, and where both the first and second chimeric PIVs elicit an immune response against the non-PIV pathogen (all claimed). Dwg.0/21

L6 ANSWER 8 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2001-103088 [11] WPIDS

DOC. NO. CPI:

C2001-030283

TITLE:

Isolated chimeric human-bovine respiratory

syncytial virus (RSV), useful

in an attenuated vaccine to elicits an immune response

against either or both human RSV A or

RSV B.

DERWENT CLASS:

B04 D16

INVENTOR(S):

BUCHHOLZ, U; COLLINS, P L; KREMPLE, C D; MURPHI, B R;

WHITEHEAD, S S; KREMPL, C D; MURPHY, B R (USSH) US DEPT HEALTH & HUMAN SERVICES

PATENT ASSIGNEE(S):

94

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA PG

WO 2001004335 A2 20010118 (200111) \* EN 148

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

NL OA PT SD SE SL SZ TZ UG ZW

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SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000056415 A 20010130 (200127)

BR 2000013195 A 20020723 (200257)

EP 1287152 A2 20030305 (200319) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

KR 2002092343 A 20021211 (200328)

CN 1402792 A 20030312 (200339)

#### APPLICATION DETAILS:

PATENT NO KI	ND .	API	PLICATION	DATE
WO 2001004335 AU 2000056415			2000-US17755 2000-56415	20000623
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EP 1287152	A2	EP	2000-US17755 2000-941756	20000623 20000624
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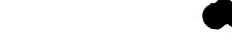
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EP 1287152		_		•		200104335

PRIORITY APPLN. INFO: US 1999-143132P 19990709

AN 2001-103088 [11] WPIDS

AB WO 200104335 A UPAB: 20030214

#### Lucas 10/073,065



NOVELTY - An isolated chimeric human-bovine respiratory syncytial virus (RSV) that is infectious and attenuated in humans, is new.

DETAILED DESCRIPTION - An isolated chimeric human-bovine respiratory syncytial virus (RSV) that is infectious and attenuated in humans, is new.

The virus comprises a major nucleocapsid protein (N), a nucleocapsid phosphoprotein (P), a large polymerase protein (L), a RNA polymerase elongation factor, and a partial or complete RSV background genome, or antigenome of a human RSV or bovine RSV, combined with one or more heterologous gene(s) or genome segment(s) of a different RSV to form a human-bovine chimeric RSV genome or antigenome.

INDEPENDENT CLAIMS are also included for the following:

- (1) a method (M1) for stimulating the immune system of an individual to induce protection against RSV, comprising administering an immunologically sufficient amount of the chimeric RSV;
- (2) an isolated polynucleotide comprising a chimeric RSV genome or antigenome which includes a partial or complete RSV background genome or antigenome of a human or bovine RSV combined with one or more heterologous gene(s) or genome segment(s) of a different RSV to form a human-bovine chimeric RSV genome or antigenome; and
- (3) a method (M2) for producing an infectious attenuated chimeric RSV particle from one or more isolated polynucleotide molecules encoding the RSV, comprising expressing RSV N, P, L and RNA polymerase elongation factor proteins, and an expression vector comprising the polynucleotide of (2) in a cell or cell-free lysate.

ACTIVITY - Antiviral.

Young chimpanzees which were determined to be seronegative for human RSV were inoculated by both the intranasal and intratracheal routes with a dose of 107 pfu (plaque forming units) per ml of rBRSV or rBRSV/A2 at each site. Each virus was administered to two chimpanzees. Following inoculation of the virus, nasopharyngeal swab samples were taken daily on days 1-10 and 12, and tracheal lavage samples were taken on days 2, 5, 6, 8 and 12. Specimens were frozen and RSV titers were measured later by plaque assay on HEp-2 cells. The amount of rhinorrhea, a measure of upper respiratory tract illness, was estimated daily and assigned a score of 0-4 (0=none, 1= trace, 2= mild, 3= moderate, 4= severe). The results were compared to historic controls of animals which had received:

- (i) 104 pfu of recombinant human RSV strain A2 wild type virus per site (Whitehead, et al., J. Virol. 72:4467-4471, 1998) or
- (ii) 105 pfu of the live-attenuated rA2cp28/404 strain A2 vaccine candidate per site (Whitehead, et al., J. Virol. 73:343 8-3442, 1999), administered by the same routes.

Wild type human RSV was highly permissive in seronegative chimpanzees, and in this exercise replicated to peak mean titers of more than 4.5 log10 pfu per mI of nasal swab or tracheal lavage sample. The peak rhinorrhea score was 2.5. The live- attenuated vaccine candidate rA2cp248/404 (see, e.g., U.S. Patent No. 5,993,824, issued November 30, 1999; International Publication No. WO 98102530; Collins, et al., Proc Natl. Acad. Sci. USA 92:11563-11567,1995; Whitehead, et al., Virology 247:232-239, 1998) replicated to mean peak titers of 2.5 and 1.4 log10 pfu per mI of swab/lavage in the upper and lower respiratory tracts, respectively, and had a peak rhinorrhea score of 0.8. In contrast, there was no detectable replication of recombinant bovine (rBRSV) in either the upper or lower respiratory tracts and no evidence of disease. Thus, even when administered at 100-1000 times the dose of human RSV, rBRSV

was highly restricted for replication in chimpanzees. The rBRSV/A2 chimera exhibited replication over several days in both the upper and lower respiratory tract.

The shedding was not detected until day 3 or 5 indicates that it was not carryover from the inoculation, as does the length of time over which virus was recovered. The titers were much lower than observed for wild type human RSV and moderately lower than observed for the rA2cp248/404 vaccine candidate. These results indicate that the chimeric virus was highly attenuated. Thus, replacement of the G and F glycoprotein genes of rBRSV with their human RSV counterparts, which transferred the major antigenic determinants, confers improved growth in chimpanzees while other bovine RSV genes contribute to a highly attenuated phenotype.

MECHANISM OF ACTION - Immunostimulant; Anti-RSV vaccine.

USE - The chimeric RSV is useful in an attenuated vaccine to elicits an immune response against either or both human RSV A or RSV B (claimed).

Dwg.0/13

L6 ANSWER 9 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2001-103086 [11] WPIDS

DOC. NO. CPI:

C2001-030281

TITLE:

Isolated infectious recombinant respiratory

syncytial virus (RSV) has a

modified genome and is used as a noninfectious subunit vaccine and for the production of viral proteins in cell

culture.

DERWENT CLASS:

B04 D16

INVENTOR(S):
PATENT ASSIGNEE(S):

BERMINGHAM, A; COLLINS, P L; MURPHY, B R (USSH) US DEPT HEALTH & HUMAN SERVICES

COUNTRY COUNT:

94

PATENT INFORMATION:

PATENT NO	O KIND	DATE	WEEK	LA	PG

WO 2001004321 A1 20010118 (200111) \* EN 124

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

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AU 2000059181 A 20010130 (200127)

#### APPLICATION DETAILS:

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WO 2001004321			2000-US18534	
AU 2000059181	A	ΑU	2000-59181	20000707

#### FILING DETAILS:

PATENT NO	KIND	•	PAT	ENT NO	
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ΔII 200005918	R1 Δ	Rased on	WO	200104321	

PRIORITY APPLN. INFO: US 1999-143097P 19990709

AN 2001-103086 [11] WPIDS AB WO 200104321 A UPAB: 20010224



NOVELTY - Isolated infectious recombinant respiratory syncytial virus (RSV) (I) comprises an RSV genome or antigenome, major nucleocapsid (N) protein, nucleocapsid phosphoprotein (P), large polymerase protein (L) and RNA polymerase elongation factor and has a modification in the genome/antigenome of the second translational open reading frame encoded by the M2 gene (M2 ORF2).

DETAILED DESCRIPTION - Isolated infectious recombinant respiratory syncytial virus (RSV)

- (I) comprises an RSV genome or antigenome, major nucleocapsid (N) protein, nucleocapsid phosphoprotein (P), large polymerase protein (L) and RNA polymerase elongation factor and has a modification in the genome/antigenome which is complete or partial deletion of the second translational open reading frame encoded by the M2 gene (M2 ORF2) or at least one nucleotide change to reduce/ablate M2 ORF2 expression.
  - INDEPENDENT CLAIMS are also included for the following:
- (1) an isolated polynucleotide molecule (II) comprising a RSV genome or antigenome modified by a partial or complete deletion of M2 ORF2 or one or more nucleotide changes that reduce or ablate expression of M2 ORF;
- (2) a method for producing an infectious attenuated RSV particle from one or more isolated polynucleotide molecules encoding the RSV;
- (3) an isolated infectious recombinant RSV (III) comprising an RSV genome or antigenome, major nucleocapsid (N) protein, nucleocapsid phosphoprotein (P), large polymerase protein (L) and RNA polymerase elongation factor and an amino acid substitution at Asn43 of the RSV polymerase gene L; and
- (4) a method for producing one or more purified RSV proteins comprising infecting a host cell permissive of RSV infection with a recombinant RSV that has an M2 ORF deletion or knock-out mutation in its genome or antigenome, isolating the recombinant RSV from the host cell and purifying the one or more RSV proteins.

ACTIVITY - Immunostimulant; respiratory general; antiinflammatory. MECHANISM OF ACTION - Vaccine; gene therapy.

Recombinant RSV unable to express NS 1 (rA2 Delta NS1) or M2-2 (rA2 Delta M2-2) viruses were administered individually to juvenile RSV-seronegative chimpanzees by combined intranasal and intratracheal inoculation at 105 pfu per ml per site. Nasopharyngeal swabs and tracheal lavage samples were collected at intervals over 10 days post infection and assayed for virus titer to monitor virus replication. The mean peak titer for the nasopharyngeal swab was 5 for wild type RSV, 1.6 for rA2 Delta NS1 and 1.5 for rA2 Delta M2-2, for the tracheal lavage the mean peak titer was 5.5 for wild type RSV, 1.2 for rA2 Delta NS1 and less than 0.7 for rA2 Delta M2-2. The chimpanzees were monitored daily for rhinorrhea, a symptom of upper. respiratory tract illness and the mean peak score determined for each group. For the wild type RSV the score was 3 (moderate), for rA2 Delta NS1 it was 2 (mild) and for rA2 Delta M2-2 it was 1.8.

USE - (I) elicits a protective immune response to RSV in a vaccinated host (claimed). This immune response is protective against serious lower respiratory tract disease e.g. pneumonia and bronchioloitis when the individual is subsequently infected with wild type RSV. (I) is administered to an individual seronegative for antibodies to RSV or possessing transplacentally acquired maternal antibodies to RSV. (I) elicits an immune response

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used for the production of viral proteins in cell culture. The M2 ORF2 deletion or knockout mutant is also used as a vector for transient gene therapy of the respiratory tract. The vector incorporates a sequence encoding a product of interest e.g. cytokines such as interleukin 2 (IL-2), IL-4, interferon gamma (IF-gamma)

and granulocyte-macrophage colony stimulating factor (GM-CSF).

ADVANTAGE - Previously a chemotherapeutic agent ribavirin and pooled donor IgG has been used to treat human RSV but these methods lack long-term effectiveness and are inappropriate for widespread use. Dwg.0/6

ANSWER 10 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN L6

against human RSV A and/or RSV B (claimed). (I) can be

ACCESSION NUMBER:

2001-081053 [09] WPIDS

DOC. NO. CPI:

C2001-023408

TITLE:

Isolated human-bovine chimeric parainfluenza virus (PIV), useful in an attenuated vaccine to elicits an immune response against one or more virus(es) selected from

human PIV1 (HPIV1), HPIV2 and HPIV3.

DERWENT CLASS:

B04 D16 INVENTOR(S):

BAILLY, J E; COLLINS, P L; DURBIN, A P; MURPHY, B R;

SCHMIDT, A C; SKIADOPOULOS, M H

PATENT ASSIGNEE(S):

(USSH) US DEPT HEALTH & HUMAN SERVICES

COUNTRY COUNT:

94

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA PG

WO 2001004320 A1 20010118 (200109)\* EN 148

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

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AU 2000056303 A 20010130 (200127)

A1 20020410 (200232) EP 1194564 ΕN

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BR 2000013190 A 20020716 (200255)

KR 2002022768 A 20020327 (200264)

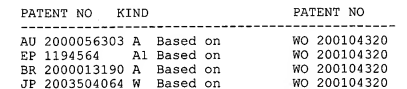
CN 1369011 A 20020911 (200282)

JP 2003504064 W 20030204 (200320) 170

#### APPLICATION DETAILS:

PAT	CENT NO K	IND	API	PLICATION	DATE
WO	2001004320	A1	WO	2000-US17066	20000616
ΑU	2000056303	A	ΑU	2000-56303	20000616
ΕP	1194564	A1	ΕP	2000-941614	20000616
			WO	2000-US17066	20000616
BR	2000013190	A	BR	2000-13190	20000615
			WO	2000-US17066	20000615
KR	2002022768	A	KR	2002-700325	20020109
CN	1369011	A	CN	2000-810120	20000616
JΡ	2003504064	W	WO	2000-US17066	20000616
		·	JΡ	2001-509524	20000616

#### FILING DETAILS:



PRIORITY APPLN. INFO: US 1999-143134P 19990709

AN 2001-081053 [09] WPIDS

AB WO 200104320 A UPAB: 20021105

NOVELTY - An isolated human-bovine chimeric parainfluenza virus (PIV) that is infectious and attenuated in humans, is new.

DETAILED DESCRIPTION - An isolated human-bovine chimeric parainfluenza virus (PIV) that is infectious and attenuated in humans, is new.

The virus comprises a major nucleocapsid protein (N), a nucleocapsid phosphoprotein (P), a large polymerase protein (L), and a partial or complete PIV background genome, or antigenome of a human PIV (HPIV) or bovine PIV (BPIV), combined with one or more heterologous gene(s) or genome segment(s) of a different PIV to form a human-bovine chimeric PIV genome or antigenome.

INDEPENDENT CLAIMS are also included for the following:

- (1) a method for stimulating the immune system of an individual to induce protection against PIV, comprising administering an immunologically sufficient amount of the chimeric PIV;
- (2) an isolated polynucleotide comprising a chimeric PIV genome or antigenome which includes a partial or complete PIV background genome or antigenome of a human or bovine PIV combined with a heterologous gene or genome segment of a different PIV to form a human-bovine chimeric PIV genome or antigenome;
- (3) a method for producing an infectious attenuated chimeric PIV particle from one or more isolated polynucleotide molecules encoding the PIV, comprising expressing PIV N, P, and L proteins, and an expression vector comprising the polynucleotide of (2) in a cell or cell-free lysate; and
- (4) an expression vector comprising an operably linked transcriptional promoter, the polynucleotide sequence of (2) and a transcriptional terminator.

ACTIVITY - Antiviral.

The rJS (wild-type HPIV3), Ka parent (Kansas BPIV3 strain), cKa (chimeric Ka strain), SF parent (Shipping fever BPIV3 strain) and cSF (chimeric SF strain) were administered intranasally and intratracheally at a dose of 100000 TCID50 per site to rhesus monkeys. Replication was monitored using standard procedures for obtaining samples from the upper (nasopharyngeal swab specimens) and lower (tracheal lavage specimens) respiratory tract and for titering the virus in LLC-MK2 cells. The cKa and cSF recombinants were significantly attenuated for the upper respiratory tract exhibiting, respectively, a 63-fold or a 32-fold reduction in mean peak virus titer compared to that of the rJS HPIV3 parent. Both cKa and cSF were also attenuated for the lower respiratory tract, but this difference was only statistically significant for cSF. The low level of replication of rJS in the lower respiratory tract made it difficult to demonstrate in a statistically-significant fashion further restriction of replication due to an attenuation phenotype at this site.

The level of replication of each chimeric virus, cKa and cSF, was not significantly different from its bovine parent in the upper or the lower respiratory tract, although the chimeric viruses each replicated better than their BPIV3 parents in the upper respiratory tract.

MECHANISM OF ACTION - Anti-PIV vaccine.

#### Lucas 10/073,065

USE - The chimeric PIV is useful in an attenuated vaccine to elicits an immune response against one or more virus(es) selected from HPIV1, HPIV2 and HPIV3.

Preferably, the chimeric PIV elicits an immune response against HPIV3 and another virus selected from HPIV1, HPIV2 or HPIV3 (claimed). Dwg.0/11

L6

ANSWER 11 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2001-091926 [10] WPIDS

DOC. NO. CPI:

C2001-027208

TITLE:

Recombinant respiratory syncytial

virus (RSV) incorporating a

heterologous polynucleotide encoding an immune modulatory

molecule is used as a vaccine to provide an immune

response to RSV.

DERWENT CLASS:

COUNTRY COUNT:

B04 D16

INVENTOR(S):

BURKREYEV, A; COLLINS, P L; MURPHY, B R; WHITEHEAD, S S;

BUKREYEV, A

PATENT ASSIGNEE(S):

(USSH) US DEPT HEALTH & HUMAN SERVICES

95

PATENT INFORMATION:

PATENT NO	KIND	DATE	WEEK	LA	PG

WO 2001004271 A2 20010118 (200110) \* EN 154

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM' DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE

SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000062112 A 20010130 (200127)

EP 1194581 A2 20020410 (200232) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

20020924 (200272) BR 2000013202 A

Α 20021211 (200324)

KR 2002092889 A 20021212 (200328)

JP 2003512817 W 20030408 (200333) 180

#### APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2001004271 A2	WO 2000-US19042	20000712
AU 2000062112 A	AU 2000-62112	20000712
EP 1194581 A2	EP 2000-948641	20000712
•	WO 2000-US19042	20000712
BR 2000013202 A	BR 2000-13202	20000712
	WO 2000-US19042	20000712
CN 1384883 A	CN 2000-810303	20000712
KR 2002092889 A	KR 2002-700505	20020114
JP 2003512817 W	WO 2000-US19042	20000712
	JP 2001-509475	20000712

#### FILING DETAILS:

PATENT NO	KIND			PAT	CENT	NO	
Att 20000621	12 A	Based	on	WO	2001	04271	

Lucas 10/073,065

EP 1194581 A2 Based on WO 200104271 BR 2000013202 A Based on WO 200104271 JP 2003512817 W Based on WO 200104271

PRIORITY APPLN. INFO: US 1999-143425P 19990713

AN 2001-091926 [10] WPIDS

AB WO 200104271 A UPAB: 20010220

NOVELTY - Infectious recombinant respiratory syncytial virus (RSV) (I) comprising a recombinant RSV genome or antigenome incorporating a heterologous polynucleotide encoding an immune modulatory molecule, a major nucleocapsid (N) protein, nucleocapsid phosphoprotein (P), large polymerase protein (L) and a RNA polymerase elongation factor, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polynucleotide molecule (II) comprising a RSV genome or antigenome modified to incorporate a polynucleotide sequence encoding an immune modulatory molecule; and
- (2) a method for producing an infectious attenuated RSV particle from one or more isolated polynucleotide molecules encoding the RSV.

ACTIVITY - Immunostimulator.

Balb/c mice were infected intranasally with 106 plaque forming units (pfu) rRSV/mIFN gamma , rRSV/chloramphenicol acetyl transferase (CAT) or wt RSV. Serum samples were collected on days 0, 28 and 56 and analyzed by RSV-specific and antibody isotype-specific enzyme linked immunosorbent assay and by an RSV neutralization assay. The levels of IgA antibodies induced by the viruses were not significantly different, there was a significant increase, four fold, in total IgG specific to RSV F protein in mice vaccinated with rRSV/mIFN gamma compared to animals vaccinated with wt RSV or RSV/CAT on day 56 but not on day 28. Neutralizing antibody titers of mice infected with rRSV/mIFN gamma compared with wt RSV and RSV/CAT were lower on day 28 but modestly higher on day 56.

MECHANISM OF ACTION - Vaccine.

USE - (I) elicits a protective immune response to RSV in a vaccinated host (claimed). (I) is administered to an individual seronegative for antibodies to RSV or possessing transplacentally acquired maternal antibodies to RSV. (I) elicits an immune response against human RSV A and/or

ADVANTAGE - (I) induces titers of serum Immunoglobulin G (IgG) that are at least 2-10 fold higher than levels of serum IgG induced by wt RSV.

Previously a chemotherapeutic agent ribavirin and pooled donor IgG has been used to treat RSV but these methods lack long-term effectiveness and are inappropriate for widespread use. Dwg.0/7

L6 ANSWER 12 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2001-071448 [08] WPIDS

CROSS REFERENCE:

2002-520377 [56]

DOC. NO. CPI:

C2001-020057

TITLE:

Obtaining an attenuated vaccine comprising recombining nucleic acids that comprise a complete or partial genomic library of a virus or cell and screening to identify those that are attenuated, useful for treating viral

infections.

DERWENT CLASS:

B04 D16

INVENTOR(S):

APT, D; DELCARDAYRE, S; HOWARD, R; PUNNONEN, J; STEMMER,





WPC

PATENT ASSIGNEE(S): COUNTRY COUNT:

(MAXY-N) MAXYGEN INC

95

PATENT INFORMATION:

PAT	CENT	NO	KIND	DATE	WEEK	LA	PG
TATO	2001	100023	3/ 7/2	20010	1104 /20010	181 * EN	117

WO 2001000234 A2 20010104 (200108)\* EN 11/

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE

SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000058809 A 20010131 (200124)

EP 1196552 A2 20020417 (200233) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

JP. 2003503039 W 20030128 (200309) 149

#### APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE .
WO 2001000234 AU 2000058809 EP 1196552		AU	2000-US16984 2000-58809 2000-944760	20000620 20000620 20000620
JP 2003503039		WO	2000-US16984 2000-US16984 2001-505941	20000620 20000620 20000620

#### FILING DETAILS:

PATENT NO K	IND	PATENT NO
AU 2000058809 EP 1196552 JP 2003503039	A2 Based on	WO 200100234 WO 200100234 WO 200100234

PRIORITY APPLN. INFO: US 1999-344655 19990625

AN 2001-071448 [08] WPIDS

CR 2002-520377 [56]

AB WO 200100234 A UPAB: 20030206

NOVELTY - A method (M1) for obtaining an attenuated vaccine comprising recombining first nucleic acids that comprise a complete or partial genomic library of a virus or cell with a second set and screening to identify those that are attenuated, is new.

DETAILED DESCRIPTION - A method (M1) of obtaining an attenuated vaccine comprises:

- (a) recombining a first set of nucleic acid segments that comprises a complete or partial genomic library of a cell with a second set of nucleic acid segments to form a library of recombinant nucleic acid fragments;
- (b) screening viruses or cells that contain members of the library of recombinant nucleic acid fragments to identify those viruses or cells that are attenuated under physiological conditions that exist in a host organism; and
- (c) screening the attenuated viruses or cells to identify those that can induce an immune response against a pathogenic agent that displays an immunogenic determinant that is also displayed by the attenuated viruses or cells.



(1) an attenuated virus or cell obtained by M1;

- (2) a vaccine composition comprising the virus or cell of (1);
- (3) a method (M2) for vaccinating an animal comprising administering the composition of (2);
- (4) a method (M3) for obtaining a chimeric attenuated vaccine comprising:

INDEPENDENT CLAIMS are also included for the following:

- (a) recombining a first set of one or more nucleic acid segments from a virus or cell with a second set of one or more nucleic acid segments, where the nucleic acid segments of the second set confer upon viruses or cells that contain the nucleic acid segments a property that is desirable for vaccination, to form a library of recombinant DNA fragments;
- (b) identifying attenuated viruses or cells by screening viruses or cells that contain members of the library of recombinant DNA fragments to identify those viruses or cells that are attenuated under physiological conditions present in a host organism inoculated with the viruses or cells; and
- (c) screening the attenuated viruses or cells to identify those that exhibit an improvement in the property that is desirable for vaccination;
- (5) a chimeric attenuated vaccine that comprises an attenuated virus or cell obtained by M3;
- (6) a vaccine composition comprising an attenuated virus or cell of
- (7) a method (M4) of vaccinating an animal comprising administering the composition of (6).

ACTIVITY - Antiviral; antibacterial; antiparasitic.

No biological data is given.

MECHANISM OF ACTION - Vaccine.

No biological data is given.

USE - The methods are useful for producing engineered attenuated vaccines which can be used against pathogenic agents such as viruses, bacteria, and parasites.

ADVANTAGE - The vaccines have improved expression of an immunogenic polypeptide, improved specific uptake, enhanced stability and enhanced immunogenecity. Dwg.0/6

ANSWER 13 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER:

2000-687044 [67] WPIDS

DOC. NO. CPI:

C2000-208979

TITLE:

Producing attenuated negative stranded RNA virus vaccines from cloned sequences, useful for immunizing against e.

q. respiratory syncytial

virus, human parainfluenza virus, Sendai virus

Newcastle disease virus, mumps virus and measles virus.

B04 C06 D16

DERWENT CLASS: INVENTOR(S):

COLLINS, P L; DURBIN, A P; MURPHY, B R; SKIADOPOULOS, M H

(USSH) US DEPT HEALTH & HUMAN SERVICES

PATENT ASSIGNEE(S): COUNTRY COUNT:

93

PATENT INFORMATION:

PATENT N	O KIND	DATE	WEEK	LA	PG
				<b>-</b>	

WO 2000061737 A2 20001019 (200067)\* EN 136

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW



AU 2000042315 A 20001114 (200108)

EP 1171623 A2 20020116 (200207) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

155

CN 1347458 A 20020501 (200252)

KR 2002008831 A 20020131 (200254)

BR 2000011159 A 20020723 (200257)

JP 2002541798 W 20021210 (200301)

#### APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE
WO 2000061737 AU 2000042315 EP 1171623		AU EP	2000-US9695 2000-42315 2000-922075 2000-US9695	20000412 20000412 20000412 20000412
CN 1347458 KR 2002008831		CN KR	2000-806224 2001-713102	20000412 20011013
BR 2000011159 JP 2002541798		WO	2000-11159 2000-US9695 2000-611661	20000412 20000412 20000412
UF 2002541790	**		2000-US9695	20000412

#### FILING DETAILS:

PATENT NO K	IND	· PA	TENT NO
AU 2000042315 EP 1171623 BR 2000011159 JP 2002541798	A2 Based A Based	on WO	200061737 200061737 200061737 200061737

PRIORITY APPLN. INFO: US 1999-129006P 19990413

AN 2000-687044 [67] WPIDS

AB WO 200061737 A UPAB: 20001223

NOVELTY - A method for producing attenuated negative stranded RNA virus vaccines from cloned sequences, is new.

DETAILED DESCRIPTION - A method (I) for producing an isolated, attenuated, recombinant negative stranded RNA virus (nsRV) from 1 or more isolated polynucleotide molecules encoding the nsRV, comprising co-expressing (in a cell or cell-free system) 1 or more expression vectors which comprise 1 or more polynucleotide molecules encoding a recombinant genome or antigenome and essential viral proteins necessary to produce an infective virus particle of the nsRV. The recombinant genome or antigenome is modified to encode a mutation within a recombinant protein of the recombinant virus at an amino acid position corresponding to an amino acid position of an attenuating mutation identified in a heterologous, mutant nsRV. The mutation, by incorporation within the recombinant protein confers an attenuated phenotype on the recombinant virus.

INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated attenuated recombinant nsRV comprising a recombinant genome or antigenome and essential viral proteins necessary to produce an infectious particle of the recombinant nsRV (the recombinant genome or antigenome is modified to encode a mutation within a recombinant protein of the virus at an amino acid position corresponding to an amino acid position of an attenuating mutation identified in a heterologous, mutant nsRV (the mutation, by incorporation within the recombinant protein confers an attenuated phenotype on the recombinant virus); and
  - (2) an expression vector comprising an operably linked

#### Lucas 10/073,065

transcriptional promoter, a polynucleotide molecule encoding a recombinant genome or antigenome of a recombinant nsRV and a transcriptional terminator (the recombinant genome or antigenome is modified to encode a mutation within a recombinant protein of the virus at an amino acid position corresponding to an amino acid position of an attenuating mutation identified in a heterologous mutant nsRV; the mutation by incorporation within the recombinant protein confers an attenuated phenotype on the recombinant virus).

ACTIVITY - Antiviral. No biological data given. MECHANISM OF ACTION - Vaccine.

USE - The recombinant viruses produced may be used for stimulating a patients immune system to induce protection against a negative stranded RNA virus (nsRV) (claimed) such as respiratory syncytial virus (RSV), especially human RSV subgroups A and B, bovine RSV, murine RSV or avian pneumovirus, human parainfluenza virus (HPIV) 1, HPIV2, HPIV 3, bovine PIV (BPIV), Sendai virus (SeV), Newcastle disease virus (NDV), simian virus 5 (SV5), mumps virus (MuV), measles virus (MeV), canine distemper virus (CDV), rabies virus (RaV) or vesicular stomatitis virus (VSV).

L6 ANSWER 14 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER: 2000

2000-679462 [66] WPIDS

DOC. NO. CPI:

Dwg.0/5

C2000-206611

TITLE:

Infectious chimeric respiratory

syncytial virus (RSV)

produced from cloned nucleotide sequences, useful as a vaccine against diseases caused by the virus, such as

pneumoniae and bronchiolitis.

DERWENT CLASS:

B04 D16

INVENTOR(S):
PATENT ASSIGNEE(S):

COLLINS, P L; MURPHY, B R; WHITEHEAD, S S (USSH) US DEPT HEALTH & HUMAN SERVICES

COUNTRY COUNT:

92

PATENT INFORMATION:

PATENT	NO	KIND	DATE	WEEK	LA	PG

WO 2000061611 A2 20001019 (200066)\* EN 278

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL

OA PT SD SE SL SZ TZ UG ZW W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ EE

ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000040655 A 20001114 (200108)

EP 1169457 A2 20020109 (200205) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

KR 2002013526 A 20020220 (200257)

BR 2000011160 A 20021008 (200277)

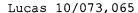
CN 1364195 A 20020814 (200280)

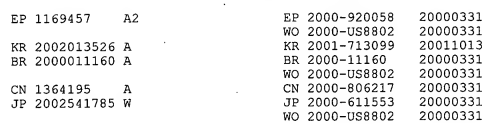
JP 2002541785 W 20021210 (200301) 309

#### APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 20000616	511 A2	WO 2000-US8802	20000331
AU 20000406		AU 2000-40655	20000331

25/08/2003





#### FILING DETAILS:

PATENT NO	KIND			PAT	ENT NO
77. 000004	0655 P	D		MO	200061611
AU 200004	0655 A .	Based	on		
EP 116945	7 A2	Based	on	WO	200061611
BR 200001	1160 A	Based	on	WO	200061611
JP 200254	1785 W	Based	on	WO	200061611

PRIORITY APPLN. INFO: US 1999-291894 19990413

AN 2000-679462 [66] WPIDS

AB WO 200061611 A UPAB: 20001219

NOVELTY - An isolated infectious chimeric respiratory

syncytial virus (RSV) comprising a major

nucleocapsid (N) protein, a nucleocapsid phosphoprotein (P), a large polymerase protein (L), an RNA polymerase elongation factor, and a partial or complete RSV genome or antigenome of one RSV strain or subgroup virus combined with a heterologous gene of a different RSV strain or subgroup virus, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) a method for stimulating the immune system of an individual to induce protection against RSV comprising administering the chimeric RSV;
- (2) an isolated polynucleotide molecule comprising a chimeric RSV genome or antigenome which includes a partial or complete RSV genome or antigenome of one RSV strain or subgroup virus combined with a heterologous gene or gene segment of a different RSV strain or subgroup virus; and
- (3) a method for producing an infectious attenuated chimeric particle from one or more isolated polynucleotide molecules encoding the RSV, comprising expressing in a cell or cell-free lysate, an expression vector comprising an isolated polynucleotide comprising a chimeric RSV genome or antigenome and RSV N, P

, L and RNA polymerase elongation factor proteins.

ACTIVITY - Antiviral.

No relevant biological data is given.

MECHANISM OF ACTION - Vaccine.

No relevant biological data is given.

USE - The chimeric respiratory syncytial

virus (RSV) is useful as a vaccine against RSV

which causes diseases such as pneumoniae and bronchiolitis in infants. Dwg.0/27

L6 ANSWER 15 OF 19 MEDLINE ON STN

DUPLICATE 2

ACCESSION NUMBER: DOCUMENT NUMBER:

2000473581 MEDLINE

20438131 PubMed ID: 10982380

TITLE:

Recombinant respiratory syncytial virus that does not express the NS1 or M2-2 protein is highly attenuated and

immunogenic in chimpanzees.

25/08/2003

AUTHOR: Teng M N; Whitehead S S; Bermingham A; St Claire M; Elkins

W R; Murphy B R; Collins P L

CORPORATE SOURCE: Respiratory Viruses Section, Laboratory of Infectious

Diseases, National Institute of Allergy and Infectious

Diseases, Bethesda, Maryland, 20892, USA.

CONTRACT NUMBER: AI-000087 (NIAID)

AI-000099 (NIAID)

SOURCE: JOURNAL OF VIROLOGY, (2000 Oct) 74 (19) 9317-21.

Journal code: 0113724. ISSN: 0022-538X.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200010

ENTRY DATE: Entered STN: 20001012

Last Updated on STN: 20001012 Entered Medline: 20001004

AB Mutant recombinant respiratory syncytial

viruses (RSV) which cannot express the NS1 and
M2-2 proteins, designated rA2DeltaNS1 and rA2DeltaM2-2,

respectively, were evaluated as live-attenuated RSV vaccines.
The rA2DeltaNS1 virus contains a large deletion that should have the

advantageous property of genetic stability during replication in vitro and in vivo. In vitro, rA2DeltaNS1 replicated approximately 10-fold less well

than wild-type recombinant RSV (rA2), while rA2DeltaM2-2 had

delayed growth kinetics but reached a final titer similar to that of rA2.

Each virus was administered to the respiratory tracts of RSV -seronegative chimpanzees to assess replication, immunogenicity,

and protective efficacy. The rA2DeltaNS1 and rA2DeltaM2-2 viruses were 2,200- to 55,000-fold restricted in replication in the upper and lower

respiratory tracts but induced a level of RSV-neutralizing antibody in serum that was only slightly reduced compared to the level

induced by wild-type RSV. The replication of wild-type

RSV in immunized chimpanzees after challenge was reduced more than 10,000-fold at each site. Importantly, rA2DeltaNS1 and rA2DeltaM2-2 were 10-fold more restricted in replication in the upper respiratory tract than

was the cpts248/404 virus, a vaccine candidate that retained mild reactogenicity in the upper respiratory tracts of 1-month-old infants.

Thus, either rA2DeltaNS1 or rA2DeltaM2-2 might be appropriately attenuated for this age group, which is the major target population for an

RSV vaccine. In addition, these results show that neither

NS1 nor M2-2 is essential for RSV replication

in vivo, although each is important for efficient replication.

L6 ANSWER 16 OF 19 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

ACCESSION NUMBER: 1998-437001 [37]

DOC. NO. CPI: C1998-132762

TITLE: Ester polymers from hydroxy acids and hydroxy amino acids

- are biocompatible and biodegradable, as carrier for

bioactive materials, e.g. vaccines, proteins,

WPIDS

anti-sense oligo-nucleotide(s), drugs.

DERWENT CLASS: A23 A96 B04 D16

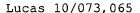
INVENTOR(S): CHONG, P; KLEIN, M H; SOKOLL, K K

PATENT ASSIGNEE(S): (CONN-N) CONNAUGHT LAB LTD; (AVET) AVENTIS PASTEUR LTD

COUNTRY COUNT: 80

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG
WO 9828357 A1 19980702 (199837) \* EN 146



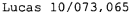


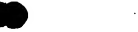
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#### APPLICATION DETAILS:

PAI	TENT NO K	IND			AP	PLICATION	DATE
	9828357	A1			MO.	1997-CA980 1998-54721 1997-951024 1997-CA980 1996-770850 1997-CA980	19971219
	9854721	Α			AU	1998-54721	19971219
EΡ	946624	A1			EP	1997-951024	19971219
					WO	1997-CA980	19971219
	6042820	A			US	1996-770850	19961220
JР	2000509428	W			WO	1997-CA980	19971219
						x > > 0 0 0 0 0 0 0 0	100,1210
BR	9714065	Α			BR	1997-14065	19971219
						1997-CA980	19971219
		A1				1999-5724	19990618
ΝZ	336718	Α				1997-336718	19971219
						1997-CA980	19971219
	729305	В				1998-54721	19971219
US	6228423	В1	Div	ex		1996-770850	19961220
						2000-501373	20000211
US	6287604	В1	Div	ex		1996-770850	19961220
						2000-502674	20000211
US	6312732	В1	Div	ex		1996-770850	19961220
						2000-499533	20000211
JP	3242118	В2				1997-CA980	19971219
	. 7.					1998-528169	19971219
JP	2002138139	A	Div	ex		1998-528169	19971219
						2001-255329	19971219
US	6471996	B1	Div	ex		1996-770850	19961220
					,	2000-499532	20000211
EΡ	946624 .	В1			EP		19971219
					_	1997-CA980	19971219
, DE	69720516	E				1997-620516	19971219
	•					1997-951024	19971219
					WO	1997-CA980	19971219







- .......

JP 1998-528169 19971219 JP 2001-255329 19971219

## FILING DETAILS:

JP 3428972

PATENT NO KIND PATENT NO							
AU 9854721 A Based on	WO 9828357						
EP 946624 Al Based on	WO 9828357						
JP 2000509428 W Based on	WO 9828357						
BR 9714065 A Based on	WO 9828357						
NZ 336718 A Based on	WO 9828357						
AU 729305 B Previous Publ.	AU 9854721						
	WO 9828357						
	US 6042820						
	US 6042820						
	US 6042820						
JP 3242118 B2 Previous Publ.	JP 200009428						
Based on	WO 9828357						
US 6471996 Bl Div ex	US 6042820						
EP 946624 B1 Based on	WO 9828357						
DE 69720516 E Based on	EP 946624						
	WO 9828357						
JP 3428972 B2 Previous Publ.	JP 2002138139						

B2 Div ex

PRIORITY APPLN. INFO: US 1996-770850 . 19961220; US 2000-501373 20000211; US 2000-502674 20000211; US 2000-499533 20000211; US 2000-499532 20000211

AN 1998-437001 [37] WPIDS

AB WO 9828357 A UPAB: 19980916

Biodegradable, biocompatible ester polymer from hydroxy acids and hydroxy (or thio) amino acids of formula (I) is new. R1-R5 = H or alkyl; R6 = H, a protecting group, a spacer molecule, or a biologically active agent; X = 0 or S; and x, y are integers. Also claimed are: (i) Preparation of the polymer comprising: (a) forming a monomer mixture containing at least one alpha -hydroxy acid and at least one pseudo amino acid having an amine protecting group with an organic solvent solution of an esterification catalyst under inert atmospheric conditions; (b) copolymerising the monomers; and (c) isolating the polymer; (ii) a particulate carrier for delivery of biologically active materials to a host comprising a polymer backbone of formula (I); (iii) a composition comprising the particulate carrier in (ii) and at least one biologically active material entrapped within; (iv) preparation of a particulate carrier for delivery of . biologically active materials to a host; (v) an immunogenic composition comprising the particulate carrier in (ii), an immunogen and a physiolgically acceptable carrier.

USE - (I) can be formed into films or microparticles, to serve as particulate carriers for slow or delayed release delivery of biologically active materials for diagnostic or therapeutic purposes. The bioactive materials are mixed into or entrapped within the copolymer, or even coupled to them, optionally through a spacer. Preferred (I) degrade in the body to benign metabolites which occur naturally, to release the bioactive agent. The bioactives are especially vaccines or similar agents which elicit an immunogenic response; examples are H, influenzae proteins, including non-proteolytic Hin-47 analogue, D15, P1, P2 and P6; influenza virus or its protein, as multivalent or monovalent influenza virus vaccine; Moraxella catarrhalis protein e.g. Tbp2 protein; and Helicobacter pylori protein, e.g. urease. Other bioactives are proteins and their mimetics, bacteria and their lysates, viruses, e.g. respiratory syncytial virus,

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virus infected cell lysates, DNA plasmids, antisense RNA, DNA, and oligonucleotides, peptides, e.g. CLTB-36 and M2, antigens, antibodies, a wide range of pharmacological agents (e.g. analgesics, antibiotics, antihypertensives, and steroids), carbohydrates, lipids, lipidated amino acids, glycolipids, haptens, or combinations of the above. Attached bioactive agents include cell bioadhesion groups, macrophage stimulators, polyamino acids, and polyethylene glycol. In diagnosis, imaging agents, together with the appropriate antibody to provide targeting, diseased tissue can be monitored or the disease identified. These can be made up as kits. Antibiotic compositions of (I) can also be used as coatings, for surgical implants, catheters, and other devices, to combat infections. Dwg.0/21

L6 ANSWER 17 OF 19 MEDLINE on STN DUPLICATE 3

ACCESSION NUMBER:

1998037604 MEDLINE

DOCUMENT NUMBER: 98037604

8037604 PubMed ID: 9371553

TITLE:

Recombinant respiratory syncytial

virus from which the entire SH gene has

been deleted grows efficiently in cell culture and exhibits site-specific attenuation in the respiratory tract of the

mouse.

AUTHOR:

Bukreyev A; Whitehead S S; Murphy B R; Collins P L

CORPORATE SOURCE:

Laboratory of Infectious Diseases, National Institute of

Allergy and Infectious Diseases, Bethesda, Maryland

20892-0720, USA.

SOURCE:

JOURNAL OF VIROLOGY, (1997 Dec) 71 (12) 8973-82.

Journal code: 0113724. ISSN: 0022-538X.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals '

ENTRY MONTH:

199712

ENTRY DATE:

Entered STN: 19980116

Last Updated on STN: 19980116 Entered Medline: 19971224

AB The small hydrophobic protein SH of human respiratory syncytial virus (RSV) is a short transmembrane surface protein of unknown function. A full-length cDNA of RSV strain A2 (subgroup A) antigenomic RNA was modified such that the entire SH gene, including the transcription signals and the complete mRNA-encoding sequence, was deleted and replaced by a synthetic intergenic region. This reduced the length of the antigenome by 398 nucleotides and ablated expression of 1 of the 10 RSV mRNAs. Recombinant virus containing this engineered deletion was recovered, and the absence of the SH gene was confirmed by reverse transcription in conjunction with Northern blot analysis of intracellular RNAs and gel electrophoresis of labeled intracellular proteins confirmed the lack of expression of the SH mRNA and protein. The absence of the SH gene did not noticeably affect RNA replication, but two effects on transcription were noted. First, synthesis of the G, F, and M2 mRNAs was increased, presumably due to their being one position closer to the promoter in the gene order. Second, transcription of genes downstream of the engineered site exhibited a steeper gradient of polarity. monolayers of HEp-2 cells, the SH-minus virus produced syncytia which were at least equivalent in size to those of the wild type and produced plaques which were 70% larger. Furthermore, the SH -minus virus grew somewhat better (up to 12.6-fold) than wild-type recombinant RSV in certain cell lines. While the function of the SH protein remains to be determined, it seems to be

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completely dispensable for growth in tissue culture and fusion function. When inoculated intranasally into mice, the SH-minus virus resembled the wild-type recombinant virus in its efficiency of replication in the lungs, whereas it replicated 10-fold less efficiently in the upper respiratory tract. In mice, the SH-minus and wild-type recombinant viruses were similarly immunogenic and effective in inducing resistance to virus challenge.

L6 ANSWER 18 OF 19 MEDLINE on STN DUPLICATE 4

ACCESSION NUMBER: 1998062142 MEDLINE

DOCUMENT NUMBER: 98062142 PubMed ID: 9400970

TITLE: Recombinant vaccinia viruses expressing the F,

G or N, but not the M2, protein of bovine

respiratory syncytial virus

(BRSV) induce resistance to BRSV challenge in the calf and

protect against the development of pneumonic lesions.

AUTHOR: Taylor G; Thomas L H; Furze J M; Cook R S; Wyld S G; Lerch

R; Hardy R; Wertz G W

CORPORATE SOURCE: Institute for Animal Health, Compton, Newbury, Berkshire,

UK.. animal.health@bbsrc.ac.uk

CONTRACT NUMBER: AI 20181 (NIAID)

SOURCE: JOURNAL OF GENERAL' VIROLOGY, (1997 Dec) 78 ( Pt 12)

3195-206.

Journal code: 0077340. ISSN: 0022-1317.

PUB. COUNTRY: ENGLAND: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199801

ENTRY DATE: Entered STN: 19980122

Last Updated on STN: 19980122 Entered Medline: 19980105

AB The immunogenicity and protective efficacy of recombinant

vaccinia viruses (rVV) encoding the F, G, N or

M2 (22K) proteins of bovine respiratory

syncytial virus (BRSV) were evaluated in calves, the natural host for BRSV. Calves were vaccinated either by scarification or intratracheally with rVV and challenged 6 to 7 weeks later with BRSV. Although replication of rVV expressing the F protein in the respiratory tract was limited after intratracheal vaccination, the levels of serum and pulmonary antibody were similar to those induced following scarification. The serum antibody response induced by the F protein was biased in favour of IgG1 antibody, whereas the G and the N proteins induced similar levels of IgG1: IgG2, and antibody was undetectable in calves primed with the M2 protein. The F protein induced neutralizing antibodies, but only low levels of complement-dependent neutralizing antibodies were induced by the G protein, and antibody induced by the N protein was not neutralizing. F and N proteins primed calves for BRSV-specific lymphocyte proliferative responses, whereas proliferative responses were detected in calves primed with the G protein only after BRSV challenge. The M2 protein primed lymphocytes in only one out of five calves. Although there were differences in the immune responses induced by the rVVs, the F, G and N, but not the M2, proteins induced significant protection against BRSV infection and, in contrast with the enhanced lung pathology seen in mice vaccinated with rVV expressing individual proteins of human (H)RSV, there was a

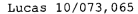
L6 ANSWER 19 OF 19 MEDLINE on STN

reduction in lung pathology in calves.

DUPLICATE 5

Page 40

25/08/2003



ACCESSION NUMBER:

1998055719 MEDLINE

DOCUMENT NUMBER:

98055719 PubMed ID: 9395341

TITLE:

Structural properties of chimeric peptides containing a T-cell epitope linked to a fusion peptide and their importance for in vivo induction of cytotoxic T-cell

responses.

AUTHOR:

Lelievre D; Hsu S C; Daubos P; Favard C; Vigny P; Trudelle

Y; Steward M W; Delmas A

CORPORATE SOURCE:

Centre de Biophysique Moleculaire, UPR 4301 CNRS, Orleans,

France.

SOURCE:

EUROPEAN JOURNAL OF BIOCHEMISTRY, (1997 Nov 1) 249 (3)

895-904.

PUB. COUNTRY: GEI

Journal code: 0107600. ISSN: 0014-2956. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)

DOCUMENT TYPE: LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199801

immunogenicity than the M2-F peptide.

ENTRY DATE:

Entered STN: 19980129

Last Updated on STN: 19980129

Entered Medline: 19980115

We have previously shown that when administered to mice without adjuvant, AΒ a chimeric peptide consisting of the fusion peptide F from measles virus protein linked at the C-terminus of a cytotoxic T-cell epitope from the M2 protein of respiratory syncytial virus efficiently primes for an major histocompatibility complex (MHC) class-I restricted cytotoxic T lymphocyte (CTL) response. In this report, we demonstrated by microspectrofluorometry that the fusion-peptide moiety bound to the plasma membrane of living cells. When the fusion peptide was linked to the C-terminus of the CTL epitope, the chimeric peptide (M2-F) adopted a marked beta-sheet conformation. In contrast, when the fusion peptide was linked to the N-terminus of the T-cell epitope ( F-M2), the chimeric peptide adopted an alpha-helical conformation in the presence of trifluoroethanol. The immunogenicity of the two chimeric peptides for class-I restricted CTL was also significantly different, the one adopting the alpha-helical conformation being more immunogenic. Probably due to its obvious conversion to an alpha-helical conformation, the F-M2 peptide could have a higher propensity to insert into

membranes, as shown by microspectrofluorometry, with a resultant better

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23/08/2003

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ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2003 ACS on STN

2003:282424 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 138:286003

DNA vaccine for respiratory syncytial virus TITLE: Mohaptra, Shyam S.; Kumar, Mukesh; Huang, INVENTOR(S):

Shau-ku; Leong, Kam W.; Lockey, Richard F.; Zhang, Jian; Behera, Aruan K.; Chen, Li-chen; Perez De

La Cruz, Ch

University of South Florida, USA; Johns Hopkins PATENT ASSIGNEE(S):

University

SOURCE: PCT Int. Appl., 35 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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KIND DATE
                                          APPLICATION NO.
     PATENT NO.
                                                           DATE
                                          -----
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                           20030410
                                         WO 2002-US4114 20020212
     WO 2003028759
                      A1
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            CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU,
            ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
            LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD,
            SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU,
             ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                           20030410
                                          US 2002-73065
                                                           20020212
     US 2003068333
                      A1
PRIORITY APPLN. INFO .:
                                       US 2001-325573P P 20010928
    An effective prophylactic mucosal gene expression
    vaccine (GXV), made up of a cocktail of a least 4 different
    plasmid DNAs encoding corresponding RSV antigens, coacervated with
     chitosan to formulate nanospheres. In a murine model of RSV infection,
     intranasal administration with GXV results in significant induction of
     RSV-specific antibodies, nasal IgA antibodies, cytotoxic T lymphocytes,
     and IFN-.gamma. prodn. in the lung and splenocytes. A single dose of GXV
     induces a drastic redn. of viral titers.
IC
     ICM A61K039-155
     ICS A61K047-36; C12N015-00
CC
     15-2 (Immunochemistry)
     Section cross-reference(s): 14, 63
ST
     respiratory syncytial virus DNA vaccine
ΙT
     Proteins
     RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
```

respiratory syncytial virus) Human respiratory syncytial virus ΙT

vaccine of plasmid vectors)

Immunoglobulins

Human

IT

IT

(DNA vaccine of plasmids expressing antigens of)

(22,000-mol.-wt., M2; DNA vaccine of plasmid vectors expressing antigens of respiratory syncytial virus)

(DNA vaccine of plasmid vectors expressing antigens of

RL: BSU (Biological study, unclassified); BIOL (Biological study) (A, secretory; to respiratory syncytial virus induced by DNA

```
ΙT
    Glycoproteins
    RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
    THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (F; DNA vaccine of plasmid vectors expressing antigens of
        respiratory syncytial virus)
    Glycoproteins
IT
     RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (G; DNA vaccine of plasmid vectors expressing antigens of
        respiratory syncytial virus)
     Immunoglobulins
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (G; to respiratory syncytial virus induced by DNA vaccine of
       plasmid vectors)
IT
    Proteins
    RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (M (matrix); DNA vaccine of plasmid vectors expressing
        antigens of respiratory syncytial virus)
     Proteins
IT
     RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (N (nucleocapsid); DNA vaccine of plasmid vectors expressing
        antigens of respiratory syncytial virus)
IT
     RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (NS1 (nonstructural, 1); DNA vaccine of plasmid vectors
        expressing antigens of respiratory syncytial virus)
IT
     Proteins
     RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (NS2 (nonstructural, 2); DNA vaccine of plasmid vectors
        expressing antigens of respiratory syncytial virus)
     Phosphoproteins
ΙT
     RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (P; DNA vaccine of plasmid vectors expressing antigens of
        respiratory syncytial virus)
     T cell (lymphocyte)
IT
        (cytotoxic; to respiratory syncytial virus induced by DNA
        vaccine of plasmid vectors)
IT
     RL: BSU (Biological study, unclassified); PAC (Pharmacological activity);
     THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (hydrophobic, SH (small hydrophobic); DNA vaccine of plasmid
        vectors expressing antigens of respiratory syncytial virus)
     Respiratory tract, disease
ΙT
        (hyperresponsiveness; DNA vaccine of plasmid vectors
        expressing antigens of respiratory syncytial virus in relation to)
     Development, mammalian postnatal
ΙT
        (infant; DNA vaccine of plasmid vectors expressing antigens
        of respiratory syncytial virus)
ΙT
     Respiratory tract, disease
        (lower, infection; plasmid vectors expressing antigens of respiratory
        syncytial virus for vaccination against)
     Drug delivery systems
IT
        (nanospheres; for plasmid vectors expressing antigens of respiratory
        syncytial virus)
     Vaccines
ΙT
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```
(nasal; plasmids expressing antigens of respiratory syncytial virus)
TΤ
    Gene, microbial
    RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (of respiratory syncytial virus antigens for DNA vaccination)
TT
    Vaccines
        (oral; plasmids expressing antigens of respiratory syncytial virus)
TΤ
    Plasmid vectors
        (pVAX; for expression of antigens of respiratory syncytial virus in DNA
        vaccination)
IT
    Interferons
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (.gamma.; DNA vaccine of plasmid vectors expressing antigens
       of respiratory syncytial virus induces prodn. of)
    9012-76-4, Chitosan
IT
    RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL
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        (nanospheres; for delivery of plasmid vectors expressing antigens of
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                        5
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REFERENCE COUNT:
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    ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2003 ACS on STN
L9
                        2002:888800 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                        137:389134
TITLE:
                        Biodegradable polyphosphates for controlled release of
                        drugs and genes and their preparation
INVENTOR(S):
                        Wang, Jun; Mao, Hai-Quan; Leong, Kam Weng
PATENT ASSIGNEE(S):
                        Johns Hopkins Singapore Pte. Ltd., Singapore
SOURCE:
                        PCT Int. Appl., 53 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
                        English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                         APPLICATION NO. DATE
    PATENT NO.
                   KIND DATE
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                    A1 20021121 WO 2002-SG90 20020514
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
            PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
            UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU,
            TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                       US 2001-290888P P 20010514
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The pos. chargeable polyphosphoester comprises .gtoreq.1 phosphoester linkage in the polymer backbone and .gtoreq.1 pos. chargeable group which is a substituent of a side chain attached to the polymer backbone through a phosphoester linkage. The polyphosphoester is prepd. by polymg. .gtoreq.1 monomer to form a polymer with .gtoreq.1 phosphoester linkage in polymer backbone, reacting the polymer with a alc. with a chargeable group or its substituents. The compns. contg. the polyphosphoesters and biol. active substances are useful for delivery of drugs and genes. A controlled gene delivery system based on these polyphosphoesters is prepd. by complex coacervation of nucleic acid (DNA or RNA) with the polymers.

The release rates can be manipulated by adjusting the charge ratios of polyphosphoesters to nucleic acids. This gene delivery system yields a higher gene expression in muscle when injected i.m. IC ICM C08G079-04 A61K047-48; A61P021-06; A61P011-06; A61P009-10; A61P001-08; ICS A61P035-00; A61P011-02; A61P001-12; A61P001-10 63-5 (Pharmaceuticals) CC polyphosphoester carrier drug gene delivery STAnimal cell line ΙT (Hek 293; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) Immunostimulants ΙT (adjuvants; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) Peptides, biological studies TT RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (amphiphilic; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) Polymers, biological studies IT RL: BUU (Biological use, unclassified); IMF (Industrial manufacture); BIOL (Biological study); PREP (Preparation); USES (Uses) (biodegradable; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) IT Drug delivery systems (carriers; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) IT Drug delivery systems (controlled-release; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) Steroids, biological studies IT RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (derivs.; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) Polyphosphoric acids IT RL: BUU (Biological use, unclassified); IMF (Industrial manufacture); BIOL (Biological study); PREP (Preparation); USES (Uses) (esters; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) IT RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (fragments; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) IT RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (plasmid; prepn. of biodegradable polyphosphates for controlled release of drugs and genes) IT Drugs Gene therapy Human Mammalia Mouse Primates (prepn. of biodegradable polyphosphates for controlled release of drugs and genes) IT Cytokines DNA Interleukin 10 Interleukin 12 Interleukin 4

Interleukin 5 Proteins RNA

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (prepn. of biodegradable polyphosphates for controlled release of drugs and genes)

- IT Therapy
  - (small mol.; prepn. of biodegradable polyphosphates for controlled release of drugs and genes)
- IT Interferons
  - RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
    (.alpha.; prepn. of biodegradable polyphosphates for controlled release
    of drugs and genes)
- IT Interferons
  - RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (.gamma.; prepn. of biodegradable polyphosphates for controlled release of drugs and genes)
- IT 83906-57-4DP, chlorinated, esterified and hydrolyzed 83945-68-0DP, chlorinated, esterified and hydrolyzed
  - RL: BUU (Biological use, unclassified); IMF (Industrial manufacture); BIOL (Biological study); PREP (Preparation); USES (Uses)
    - (prepn. of biodegradable polyphosphates for controlled release of drugs and genes)
- TT 77987-49-6DP, Benzyl N-(2-hydroxyethyl)carbamate, reaction products of chlorinated poly(4-methyl-2-hydro-1,3,2-dioxaphospholane)
  - RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)
    - (prepn. of biodegradable polyphosphates for controlled release of drugs and genes)
- IT 64-18-6, Formic acid, reactions 7782-50-5, Chlorine, reactions
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- IT 9031-11-2, .beta.-Galactosidase 191681-52-4, Sequence
  RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
  (prepn. of biodegradable polyphosphates for controlled release of drugs and genes)
- IT 16352-26-4
  - RL: RCT (Reactant); RACT (Reactant or reagent)
    (starting material; prepn. of biodegradable polyphosphates for controlled release of drugs and genes)
- REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

# => d his ful

(FILE 'HOME' ENTERED AT 15:56:41 ON 25 AUG 2003)

L1 L2 L3 L4	4360 SEA ABB=ON 187 SEA ABB=ON	CD AT 15:56:51 ON 25 AUG 2003  I (RSV? OR ?RESPIRATOR?(W)?SYNCYTIAL?(W)?VIRUS?)  I L1 AND ?IMMUNOGEN?  I L2 AND (F OR G OR M OR SH OR NS1? OR NS2? OR P)  I L3 AND M2? /3 ali from CA Plus
L5 L6	10.00 FF ON OF BUILD OF	s, EMBASE, WPIDS, JICST-EPLUS, JAPIO' ENTERED AT 2003 I L4 L5 (10 DUPLICATES REMOVED) 19 Celá from Other database

Please let me know if you'd like any revisions -Many Jackahl